FILE 'HOME' ENTERED AT 12:58:40 ON 06 MAR 2001

- => file biosis caba caplus embase lifesci medline scisearch uspatfull japio
- => e mardh sven/au
- E1 1 MARDH PERANDERS/AU
- E2 337 MARDH S/AU
- E3 39 --> MARDH SVEN/AU
- E4 6 MARDHEKAR B V/AU
- E5 1 MARDHEKAR DHANANJAY V/AU
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- E8 4 MARDHY ABDELHAKIM/AU
- E9 7 MARDI A/AU
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- E11 1 MARDI A S/AU
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- => s e2-e3 and (gastri? or pylori)
- LI 243 ("MARDH S"/AU OR "MARDH SVEN"/AU) AND (GASTRI? OR PYLORI)
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L2 86 DUP REM L1 (157 DUPLICATES REMOVED)

=> d bib ab 1-

YOU HAVE REQUESTED DATA FROM 86 ANSWERS - CONTINUE? Y/(N):y

L2 ANSWER 1 OF 86 CAPLUS COPYRIGHT 2001 ACS

AN 2001:128060 CAPLUS

- TI Characterization of oxyntic glands isolated from the rat ***gastric***
 mucosa
- AU Azerkan, L.; Bengtsson, P.; Tommeras, K.; Li, Z.-Q.; ***Mardh, S.***
- CS Faculty of Health Sciences, Division of Cell Biology, Department of Biomedicine and Surgery, Linkoping University, S-581 85, Linkoping, Swed.
- SO Comp. Biochem. Physiol., Part A: Mol. Integr. Physiol. (2001), 128(2), 349-357
 - CODEN: CBPAB5; ISSN: 1095-6433
- PB Elsevier Science Inc.
- DT Journal
- LA English
- AB A simple and reproducible method for isolating oxyntic glands from the rat

 gastric mucosa was developed. The mucosa was incubated with
 pronase and EGTA, and then treated mech. to release glands that were sepd.
 from single cells by sedimentation. Parietal cells were identified by
 immunostaining using a monoclonal antibody against H,K-ATPase. The
 glandular cells appeared morphol. intact. By careful control of the
 conditions of gland isolation, long glandular structures comprising
 hundreds of cells surrounding the lumen were obtained. I.p. injection of

Br-deoxyuridine in the rat 1.5 h before the isolation procedure resulted in glands with a labeling of cells in their neck region. The glands were viable, as demonstrated by their ability to respond to various hormones. Histamine dose-dependently stimulated the acid formation which was measured as the accumulation of [14C]aminopyrine. At 100 mu.M histamine the accumulation was increased 5-10-fold. At 100 nM, pentagastrin potentiated the histamine stimulated accumulation by approx. 40% but pentagastrin alone did not stimulate. The oxyntic glands obtained by the present procedure appear useful for studies on cell physiol., including regulation of acid secretion, cellular interactions, and possibly also differentiation and proliferation mechanisms since long glandular fragments that contained the proliferative zone could be isolated.

L2 ANSWER 2 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 1

AN 2001:25050 BIOSIS

DN PREV200100025050

- TI Omeprazole and CYP2C19 polymorphism: Effects of long-term treatment on ***gastrin***, pepsinogen I, and chromogranin A in patients with acid related disorders.
- AU Sagar, M.; Bertilsson, L.; Stridsberg, M.; Kjellin, A.; ***Mardh, S.***; Seensalu, R. (1)
- CS (1) Department of Medicine, St Gorans Sjukhus AB, S-112 81, Stockholm: Rein.Seensalu@stgoran.se Sweden
- SO Alimentary Pharmacology & Therapeutics, (November, 2000) Vol. 14, No. 11, pp. 1495-1502. print. ISSN: 0269-2813.

DT Article

LA English

SL English

AB Background: The polymorphic enzyme CYP2C19 is of importance for the metabolism and effects of omeprazole during short-term treatment. Aim: To investigate the relationship between CYP2C19 genotype and the effects of long-term omeprazole treatment. Material and methods: A total of 180 patients with acid related disorders were genotyped for wild type and mutated CYP2C19 alleles by allele-specific PCR amplification.

Gastrin and chromogranin A were assessed by radioimmunoassays, and pepsinogen I and H. ***pylori*** serology were assessed by ELISA methods. Results: In 108 of the patients, who received a single dose of 20 mg omeprazole, there was no difference in ***gastrin*** and chromogranin A concentrations between the three CYP2C19 genotypes. In 72 patients on long-term treatment (> 1 year) with 20 mg omeprazole daily. serum ***gastrin*** as well as plasma chromogranin A concentrations (mean +- s.e.) were both about threefold higher in the wild type/mutated (52.1 + -7.6 pM and 7.3 + -1.3 nM (n = 19), respectively) compared to wildtype/wild type (14.7 +- 0.9 pM and 2.5 +- 0.1 nM (n = 52), respectively; both comparisons P = 0.0001). In a single mutated/mutated patient on long-term treatment, both ***gastrin*** and chromogranin A were high (88 pM and 13.7 nM, respectively). Serum pepsinogen I concentration was significantly lower in wild type/mutated (n = 19) patients on long-term treatment, compared with the corresponding wild type/wild type (n = 49) group (147 +- 19 mug/L vs. 193 +- 12 mug/L, P = 0.04). Conclusion: Patients with one (and probably also with two) mutated CYP2C19 allele(s) on long-term treatment with omeprazole had significantly affected serum

gastrin and pepsinogen I and plasma chromogranin A concentrations compared with patients with two normal alleles. This indicates that

changes in ***gastric*** mucosal morphology during omeprazole treatment might be dependent upon the degree of the individual's capacity to metabolize omeprazole.

L2 ANSWER 3 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 2

AN 2000:406170 BIOSIS

DN PREV200000406170

- TI Prevalence of gastroduodenitis and Helicobacter ***pylori*** infection in a general population sample: Relations to symptomatology and life-style.
- AU Borch, Kurt (1); Jonsson, Kjell-Ake; Petersson, Fredrik; Redeen, Stefan; ***Mardh, Sven***; Franzen, Lennart E.
- CS (1) Department of Surgery, University Hospital, S-58185, Linkoping Sweden
- SO Digestive Diseases and Sciences, (July, 2000) Vol. 45, No. 7, pp. 1322-1329. print.

ISSN: 0163-2116.

DT Article

LA English

SL English

- AB Some benign and malignant diseases develop on the background of chronic ***gastritis*** or duodenitis. The present study was performed in order to determine the magnitude of these background changes with relations to symptomatology and life style in the general population. Examinations were performed in 501 volunteers (age 35-85 years). Fifty percent had
 - ***gastritis***; this was associated with H. ***pylori*** in 87%. H. ***pylori*** -negative ***gastritis*** was associated with regular use of NSAIDs (odds ratio 3.8 (1.6-9.9)). Duodenitis, observed in 32%, was associated with H. ***pylori*** infection (odds ratio 2.3 (1.3-4.6)), previous cholecystectomy (odds ratio 3.6 (1.1-16.1)), and regular use of NSAIDs (odds ratio 3.0 (1.4-7.1)). Neither ***gastritis*** nor duodenitis was associated with smoking or alcohol consumption. The rate of digestive symptoms did not differ between subjects with and without uncomplicated ***gastritis*** or duodenitis. In conclusion, half of this adult population had ***gastritis*** strongly associated with H.
 - ***pylori*** infection. ***Gastritis*** without H. ***pylori*** infection was frequently associated with regular NSAID intake. One third had duodenitis, which was associated with H. ***pylori*** infection as well as with regular use of NSAIDs and previous cholecystectomy. Digestive symptoms were not overrepresented in uncomplicated ***gastritis*** or duodenitis.

L2 ANSWER 4 OF 86 CAPLUS COPYRIGHT 2001 ACS

AN 2000:788838 CAPLUS

DN 134:143721

- TI N-terminal phosphorylation of ***gastric*** H/K-ATPases both in vitro and in vivo
- AU Kanagawa, M.; Umezu, H.; Kaya, S.; Watanabe, S.; Kagawa, I.; Shimada, A.; Imagawa, T.; ***Mardh, S.***; Taniguchi, K.
- CS Biological Chemistry, Division of Chemistry, Graduated School of Science, Hokkaido University, Sapporo, 060-0852, Japan
- SO Int. Congr. Ser. (2000), 1207(Na/K-ATPase and Related ATPases), 587-590 CODEN: EXMDA4; ISSN: 0531-5131
- PB Elsevier Science B.V.
- DT Journal
- LA English

AB Acid secretion is regulated by second messenger pathways. Histamine stimulates acid secretion via the H2 receptor-mediated activation of cAMP dependent protein kinase. Acetylcholine also stimulates acid secretion by increasing protein kinase C activity, via an increase in intercellular calcium concns. This protein kinase-dependent acid secretion is regarded to be coupled with intercellular protein phosphorylation. Tyr kinase modifiers regulate acid secretion. The Tyr-10 and Tyr-7 in the .alpha.-chain of not only pig but also rat and rabbit stomach H/K-ATPase of the G1 fraction are reversibly phosphorylated by endogenous Tyr kinase and phosphatase. Time course of phosphotyrosine (PY) formation in the G1 fractions from these animals with or without vanadate indicated the presence of vanadate-sensitive Tyr phosphatase in each G1 fractions. Mild tosylphenylalanyl chloromethyl ketone-trypsin treatment of the phosphorylated .alpha.-chain completely abolished PY with little detectable change in the mobility of the .alpha.-chains. These data indicate that a reversible Tyr phosphorylation occurs at N-terminal domain of the .alpha.-chain from these animals. To investigate whether the phosphorylation occurs in vivo, minced rat, rabbit and pig stomach tissues were incubated with pervanadate (PV), which permits the detection of Tyr phosphorylation through the irreversible inhibition. The results obtained indicate that a c-Src kinase present in the G1 membrane phosphorylates the N-terminal domain of the H/K-ATPase .alpha.-chain.

RE.CNT 12

RE

- (1) Chew, C; Am J Physiol 1980, V238, PG312 CAPLUS
- (2) Chew, C; Am J Physiol 1994, V267, PG818 CAPLUS
- (3) Chew, C; Biochim Biophys Acta 1986, V888, P116 CAPLUS
- (4) Chiba, T; Am J Physiol 1988, V255, PG99 CAPLUS
- (5) Hersey, S; Biochim Biophys Acta 1983, V755, P293 CAPLUS
- ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L2 ANSWER 5 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 3
- AN 2000:157078 BIOSIS
- DN PREV200000157078
- TI Expression of extracellular matrix proteins in the fetal rat ***gastric*** mucosa.
- AU Tommeras, Karin (1); Cabero, Jose Luis; ***Mardh, Sven***
- CS (1) Department of Biomedicine and Surgery, Division of Cell Biology, Faculty of Health Sciences, Linkoping University, S-581 85, Linkoping Sweden
- SO Anatomy and Embryology., (March, 2000) Vol. 201, No. 3, pp. 149-156. ISSN: 0340-2061.
- DT Article
- LA English
- SL English
- AB At gestational day 16 the epithelium of the rat stomach consists of a stratified layer of undifferentiated cells, and two days later glandular structures appear. The present study was carried out to identify extracellular matrix proteins that could be involved in the epithelial cell proliferation and differentiation processes that occur in the fetal rat stomach during this period. For comparative purposes the expression of the same components in the adult ***gastric*** mucosa was examined. Pregnant Sprague-Dawley rats received an intraperitoneal injection of 5-bromo-2'-deoxyuridine to label proliferating cells. One, 3.5, or 6 h post-injection the stomachs were excised and immediately frozen. The

specimens were sectioned and stained with hematoxylin and eosin or for 5-bromo-2'-deoxyuridine, cytokeratin no. 8, H,K-ATPase, and the extracellular matrix proteins fibronectin, laminin, and collagens type I and IV. A stratified layer of proliferating cells was observed in the epithelium of the fetal stomachs, while in adult stomachs proliferating cells were detected in the isthmus/neck region of the glands. Cytokeratin, an epithelial cell marker, was sparse at gestational day 16 but abundant both at gestational day 18 and in the isthmus/neck region of ***gastric*** glands of the adult stomach. The parietal cell marker H.K-ATPase could not be detected in the fetal stomachs during this period. Fibronectin was observed in the stroma of both fetal and adult stomachs. Collagen type I could only be detected in the stroma close to the oesophagus at gestational day 16. Two days later, collagen type I was abundant in the lamina propria, the submucosa and in the serosa of the fetal stomachs. In adult tissue collagen type I was detected in the surface epithelium, the submucosa and in the serosa of the stomach. Collagen type IV and laminin were expressed in the lamina propria, the basement membranes around blood vessels, muscle cells, and nerve bundles. as well as in the serosa of both 16- and 18-day-old fetal and adult rat stomachs. In conclusion, a high cell proliferation rate was observed in the epithelium at both gestational days 16 and 18. The increased expression of cytokeratin observed during this period indicates that the epithelial character of the embryonic cells becomes more distinct, while the remarkable change in the expression of collagen type I might reflect an important role of collagen type I in the development of the ***gastric*** epithelium.

L2 ANSWER 6 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 4

AN 2000:229681 BIOSIS

DN PREV200000229681

TI Helicobacter ***pylori*** -antigen-binding fragments expressed on the filamentous M13 phage prevent bacterial growth.

AU Cao, Jun, Sun, Yi-qian; Berglindh, Thomas; Mellgard, Bjorn; Li, Zhao-qi; Mardh, Bibbi; ***Mardh, Sven (1)***

CS (1) Department of Biomedicine and Surgery, Division of Cell Biology, Faculty of Health Sciences, Linkoping University, Linkoping Sweden

SO Biochimica et Biophysica Acta, (March 6, 2000) Vol. 1474, No. 1, pp. 107-113.

ISSN: 0006-3002.

DT Article

LA English

SL English

AB Colonization of the human stomach by Helicobacter ***pylori*** is associated with the development of ***gastritis***, duodenal ulcer, mucosa-associated lymphoid tissue (MALT) lymphoma, and ***gastric*** cancer. H. ***pylori*** -antigen-binding single-chain variable fragments (ScFv) were derived from murine hybridomas producing monoclonal antibodies and expressed as a g3p-fusion protein on a filamentous M13 phage. The recombinant ScFv-phage reacted specifically with a 30-kDa monomeric protein of a H. ***pylori*** surface antigen preparation and by means of immunofluorescence microscopy the phage was shown to bind to both the spiral and coccoid forms of the bacterium. In vitro, the recombinant phage exhibited a bacteriocidal effect and inhibited specifically the growth of all the six strains of H. ***pylori*** tested. When H. ***pylori*** was pretreated with the phage 10 min

before oral inoculation of mice, the colonization of the mouse stomachs by the bacterium was significantly reduced (P < 0.01). The results suggest that genetic engineering may be used to generate therapy-effective phages.

L2 ANSWER 7 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS **DUPLICATE 5**

AN 2000:337801 BIOSIS

DN PREV200000337801

- TI Effects of cholecystokinin on acid formation in glands and cells isolated from rabbit and rat ***gastric*** mucosa.
- AU Bengtsson, Per (1); Azerkan, Leila; Lundqvist, Gudmar; Nilsson, Goran; ***Mardh, Sven***
- CS (1) Department of Biomedicine and Surgery, Faculty of Health Sciences. Linkoping University, S-581 85, Linkoping Sweden
- SO Comparative Biochemistry and Physiology Part A Molecular & Integrative Physiology, (May, 2000) Vol. 126A, No. 1, pp. 77-84. print. ISSN: 1095-6433.

DT Article

LA English

SL English

- AB Isolated ***gastric*** glands and isolated cells prepared from rabbit and rat were studied to analyse the influence of cholecystokinin octapeptide (CCK 8) on histamine stimulated parietal cell acid formation as assessed by (14C)aminopyrine sequestered in acid tissue compartments. In rabbit ***gastric*** glands, CCK 8 evoked 32+-6% (P < 0.01) inhibition of histamine stimulated acid formation, whereas in glands prepared from rat no inhibition was recorded. Instead, CCK 8 seemed to induce a variable increase of the histamine stimulation in rat ***gastric*** glands as the aminopyrine accumulation was increased by 110+-46% (P < 0.1). Further studies on cell preparations derived from rabbit ***gastric*** mucosa revealed dual properties of CCK 8, eliciting either inhibition or stimulation of the parietal cell depending on the presence of endocrine cells. The results show that paracrine communication may be effective in glandular preparations, but seems to vary depending on species.
- L2 ANSWER 8 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 1999:445434 SCISEARCH

GA The Genuine Article (R) Number: 187GJ

TI Expression of CCK-B/ ***gastrin*** receptors in undifferentiated epithelial cells of fetal rat stomachs.

AU Tommeras K (Reprint); Cabero J L; Forssmann W G; ***Mardh S***

CS FAC HLTH SCI, DEPT BIOMED & SURG, LINKOPING, SWEDEN; ASTRA HASSLE AB, DEPT BIOCHEM & CELL BIOL, MOLNDAL, SWEDEN; LOWER SAXONY INST PEPTIDE RES, HANNOVER, GERMANY

CYA SWEDEN; GERMANY

SO GASTROENTEROLOGY, (APR 1999) Vol. 116, No. 4, Part 2, pp. G2842-G2842. Publisher: W B SAUNDERS CO, INDEPENDENCE SQUARE WEST CURTIS CENTER, STE 300, PHILADELPHIA, PA 19106-3399. ISSN: 0016-5085.

DT Conference; Journal

FS LIFE; CLIN

LA English

REC Reference Count: 0

L2 ANSWER 9 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1999:314082 BIOSIS

DN PREV199900314082

TI Expression of CCK-B/ ***gastrin*** receptors in undifferentiated epithelial cells of fetal rat stomachs.

AU Tommeras, K. (1); Cabero, J. L.; Forssmann, Wolf-Georg; ***Mardh, S.***

CS (1) Dept of Biomed and Surg, Faculty of Health Sci, Linkoping Sweden

SO Gastroenterology, (April, 1999) Vol. 116, No. 4 PART 2, pp. A652. Meeting Info.: Digestive Disease Week and the 100th Annual Meeting of the American Gastroenterological Association Orlando, Florida, USA May 16-19, 1999 American Gastroenterological Association ISSN: 0016-5085.

DT Conference

LA English

L2 ANSWER 10 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 1999:644918 SCISEARCH

GA The Genuine Article (R) Number: 226GJ

TI Direct evidence for in vivo reversible tyrosine phosphorylation of the N-terminal domain of the H/K-ATPase alpha-subunit in mammalian stomach cells

AU Kanagawa M; Kaya S; Umezu H; Watanabe S; Togawa K; Shimada A; Imagawa T; ***Mardh S***; Taniguchi K (Reprint)

CS HOKKAIDO UNIV, GRAD SCH SCI, SAPPORO, HOKKAIDO 060081, JAPAN (Reprint), HOKKAIDO UNIV, GRAD SCH SCI, SAPPORO, HOKKAIDO 060081, JAPAN, LINKOPING UNIV, FAC HLTH SCI, DEPT BIOMED & SURG, S-58185 LINKOPING, SWEDEN CYA JAPAN, SWEDEN

SO JOURNAL OF BIOCHEMISTRY, (AUG 1999) Vol. 126, No. 2, pp. 266-270. Publisher: JAPANESE BIOCHEMICAL SOC, ISHIKAWA BLDG-3F, 25-16 HONGO-5-CHOME, BUNKYO-KU, TOKYO 113, JAPAN. ISSN: 0021-924X.

DT Article; Journal

FS LIFE

LA English

REC Reference Count: 29

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB In vivo reversible phosphorylation of Tyr-7 and Tyr-10 of the pig stomach H/K-ATPase alpha-chain was initially demonstrated in mammals, rat, rabbit, and pig, in the presence of vanadate+H2O2. In vitro phosphorylation has also been unequivocally demonstrated via the use of protease inhibitors during membrane H/K-ATPase preparation. An amphoretic detergent permitted each intrinsic kinase to phosphorylate each fusion protein containing the requisite Tyr residues, along with a reduction in alpha-chain phosphorylation. These and other data suggest that some important enzyme systems are present in the apical membrane and that they are in sufficient proximity to participate in the reversible phosphorylation of the amino terminal soluble domain of the alpha-chain with an unknown physiological function in the membrane embedded H/K-ATPase.

L2 ANSWER 11 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 6

AN 1998:258345 BIOSIS

DN PREV199800258345

TI Asymptomatic Helicobacter ***pylori*** ***gastritis*** is associated with increased sucrose permeability.

AU Borch, Kurt (1); Sjostedt, Camilla; Hannestad, Ulf; Soderholm, Johan D.;

Franzen, Lennart; ***Mardh, Sven***

CS (1) Dep. of Surgery, University Hospital, S-58185 Linkoping Sweden SO Digestive Diseases and Sciences, (April, 1998) Vol. 43, No. 4, pp. 749-753.

ISSN: 0163-2116.

DT Article

LA English

AB Our aim was to investigate whether there are changes in permeability to sucrose in asymptomatic Helicobacter ***pylori*** ***gastritis*** . Nineteen asymptomatic subjects with Helicobacter ***pylori*** associated ***gastritis*** with no or mild mucosal atrophy and 19 ageand sex-matched normal controls were studied by peroral load of sucrose (100 g). The fraction of the given oral dose of sucrose excreted in urine was increased in subjects with Helicobacter ***pylori*** ***gastritis*** (median 0.08% versus 0.04% in controls). Sucrose excretion was not related to atrophy, intestinal metaplasia, or inflammation in the ***gastric*** mucosa. However, sucrose permeability was related to the degree of inflammatory (neutrophil) activity, since moderate activity was associated with higher sucrose excretion than mild activity (median 0.13% vs 0.07%). Asymptomatic Helicobacter ***pylori*** ***gastritis*** was associated with an increased sucrose permeability, which could be a sign of ***gastric*** mucosal leakage. This could have implications for the diseases and complications associated with Helicobacter ***pylori*** infection.

L2 ANSWER 12 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 7

AN 1998:405941 BIOSIS

DN PREV199800405941

- TI Identification of Helicobacter in ***gastric*** biopsies by PCR based on 16S rDNA sequences: A matter of little significance for the prediction of H. ***pylori*** -associated ***gastritis***
- AU Tiveljung, Annika; Borch, K.; Jonasson, J.; ***Mardh, S.***; Petersson, F.; Monstein, H.-J. (1)
- CS (1) Div. Clinical Microbiol., Fac. Health Sci., Univ. Linkoping, Linkoping Sweden
- SO Journal of Medical Microbiology, (Aug., 1998) Vol. 47, No. 8, pp. 695-704. ISSN: 0022-2615.

DT Article

LA English

AB The aim of the present study was to correlate molecular evidence of the presence of Helicobacter ***pylori*** in ***gastric*** biopsy samples, based on analysis of 16S rDNA, vacuolating toxin (vacA), urease A (ureA) and cagA genes, with the clinical, histological and serological findings in patients with H. ***pylori*** -associated ***gastritis*** . Fresh biopsy samples were collected from the ***gastric*** antrum and corpus of 22 asymptomatic volunteers with or without H. ***pylori*** -associated ***gastritis*** . Total DNA was extracted from the biopsy material and subjected to 16S rDNA PCR amplification, Southern blotting and 16S rDNA sequence analysis of the PCR products. The vacA, ureA and cagA genes were characterised by PCR amplification and Southern blot analysis. Based on partial 16S rDNA sequence analysis, DNA belonging to the genus Helicobacter was detected in ***gastric*** biopsy samples from 20 of 22 subjects, including seven of nine histologically and serologically normal controls. Six of 20 partial 16S rDNA sequences revealed variations within variable regions V3 and V4 that deviated from

those of the H. ***pylori*** type strain ATCC 4350T and, therefore, possibly represented other species of Helicobacter. VacA genes identical with those of the type strain were found predominantly in the subjects with H. ***pylori*** ***gastritis***, and all the patients except one were found to be cagA-positive. There was no evidence of false positive PCR reactions. In conclusion, the PCR-based molecular typing methods used here were apparently too sensitive when applied to the detection of H. ***pylori*** in human ***gastric*** tissues. The lack of quantitative analysis makes them inappropriate as clinical tools for the diagnosis of H. ***pylori*** -associated ***gastritis***, despite the fact that they provide a qualitative and sensitive tool for the detection and characterisation of H. ***pylori*** in the gastrointestinal tract.

L2 ANSWER 13 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 8

AN 1998:118862 BIOSIS

DN PREV199800118862

TI Protein kinase-dependent phosphorylation of H,K ATPase-containing membranes from rat and pig stomachs.

AU Kaya, S. (1); ***Mardh, S.***

CS (1) Biol. Chem., Dep. Chem., Grad. Sch. Sci., Hokkaido Univ., Sapporo 060 Japan

SO Acta Physiologica Scandinavica, (Jan., 1998) Vol. 162, No. 1, pp. 57-62. ISSN: 0001-6772.

DT Article

LA English

AB Previously H, K ATPase preparations from pig stomach were shown to contain intrinsic protein kinase activities which phosphorylated specific tyrosine and serine residues in the N-terminal of the alpha-chain of H,K ATPase (Togawa et at. 1996). In the present investigation, pig H.K. ATPase-containing membrane preparations were compared with rat preparations. In contrast to results obtained with the alpha-subunit of H,K ATPase from pig, phosphorylation was not observed in the rat enzyme. Addition of rat preparations to the pig preparations resulted in decreased phosphorylation in pig preparations. To follow the phosphorylation of membrane proteins in vivo, 32P-loaded ***gastric*** cells prepared from rat were stimulated with several secretagogues. Proteins with molecular weights of about 120 and 80 kDa were markedly phosphorylated upon stimulation, but the alpha-subunit of H,K ATPase was not. These results suggest that phosphorylation of tyrosine or serine residues of H,K ATPase found in pig H,K ATPase preparations may not be involved in the acid secretion pathway.

L2 ANSWER 14 OF 86 CAPLUS COPYRIGHT 2001 ACS

AN 1997:542514 CAPLUS

DN 127:186628

TI Bacteriophages presenting recombinant surface protein fusion products with bacteria antigen-binding antibody for use in bacterial infection treatment

IN ***Mardh, Sven***

PA Mardh, Sven, Swed.

SO PCT Int. Appl., 34 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

PI WO 9729185 A1 19970814 WO 1997-SE172 19970205

W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC. LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML. MR, NE, SN, TD, TG

SE 9600434 A 19970807 SE 1996-434 19960206

SE 506771 C2 19980209

CA 2244792 AA 19970814 CA 1997-2244792 19970205

AU 9716817 A1 19970828 AU 1997-16817 19970205

B2 19991118 AU 712767

EP 1997-902815 19970205 EP 889955 A1 19990113

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI

CN 1210558 A 19990310 CN 1997-192116 19970205 JP 2000505648 T2 20000516 JP 1997-528446 19970205

NO 9803456 A 19981006 NO 1998-3456 19980727

PRAI SE 1996-434 19960206 WO 1997-SE172 19970205

AB The present invention relates to bacteriophages for use in the treatment or prophylaxis of bacterial infections, esp. mucosal bacterial infections such as Helicobacter ***pylori*** infections. In particular, it relates to modified filamentous bacteriophages, e.g., M13 phages, for such use, which bacteriophages present at its surface a recombinant protein comprising: (i) a first component derived from a bacteriophage surface protein; and (ii) a second component comprising variable region sequences of an antibody to provide a bacterial antigen binding site, said second component rendering said bacteriophage capable of binding to and thereby inhibiting growth of bacterial cells involved in the etiol. of said infection.

L2 ANSWER 15 OF 86 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.DUPLICATE 9 AN 1998004376 EMBASE

- TI Phosphorylation of Tyr7, Tyr10, and Ser27 of .alpha.-chain in H+,K+-ATPase by intrinsic and extrinsic kinases.
- AU Togawa K.; Kaya S.; Mori M.; Shimada A.; Imagawa T.; Taniguchi K.; ***Mardh S.***; Corbin J.; Kikkawa U.
- CS K. Taniguchi, Biological Chemistry, Graduate School of Science, Hokkaido University, Sapporo 060, Japan. KTAN@hucc.hokudai.ac.jp
- SO Annals of the New York Academy of Sciences, (1997) 834/- (582-584). Refs: 10

ISSN: 0077-8923 CODEN: ANYAA

CY United States

DT Journal; Conference Article

FS 029 Clinical Biochemistry

LA English

L2 ANSWER 16 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 10

AN 1998:95114 BIOSIS

DN PREV199800095114

- TI Detection of spiral and coccoid forms of Helicobacter ***pylori*** using a murine monoclonal antibody.
- AU Cao, Jun; Li, Zhao Q.; Borch, Kurt; Petersson, Fredrik; ***Mardh, Sven***

 *** (1)***
- CS (1) Div. Cellbiol., Dep. Biomed. Surg., Fac. Health Sci., Linkoping Univ., S-581 85 Linkoping Sweden
- SO Clinica Chimica Acta, (Nov. 28, 1997) Vol. 267, No. 2, pp. 183-196. ISSN: 0009-8981.
- DT Article
- LA English
- AB Helicobacter ***pylori*** is the major cause of ***gastritis***. The aim of this investigation was to develop a specific antibody, which recognizes both coccoid and spiral forms of Helicobacter ***pylori*** and to test this antibody on ***gastric*** biopsy sections known to harbour coccoid bacteria. Murine monoclonal antibodies against glycine-acid extracts of five strains of Helicobacter ***pylori*** were raised. Immunofluorescence and immunoelectron microscopy showed that one antibody of the IgG1 subclass was specific for both the spiral and coccoid forms. It reacted with a 28 kDa protein that was present in all the five strains tested. Using this antibody in an indirect immunofluorescence assay of formalin-fixed antral and corpus biopsy specimens from Helicobacter ***pylori*** -associated ***gastritis*** patients showed that nine of the nine antral and five of six corpus specimens harboured the coccoid form of Helicobacter ***pylori*** This technique thus provides a rapid and specific detection of both the spiral and coccoid forms.
- L2 ANSWER 17 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 11
- AN 1997:129289 BIOSIS
- DN PREV199799421102
- TI Proliferation and differentiation of cells from explants of fetal rat stomach.
- AU Tommeras, K.; Chen, Y.; Rhedin, M.; Cabero, J. L.; ***Mardh, S. (1)***
- CS (1) Dep. Cell Biol., Fac. Health Sci., Linkoping Univ., S-581 85 Linkoping Sweden
- SO Acta Physiologica Scandinavica, (1997) Vol. 159, No. 2, pp. 155-161. ISSN: 0001-6772.
- DT Article
- LA English
- AB The current understanding of the mechanisms controlling the proliferation and differentiation of the stem cells of the ***gastric*** oxyntic glands is limited. The aim of the present study was to develop a method for investigating proliferation and differentiation of undifferentiated cells from fetal rat stomach. Outgrowth of cells was initiated from explants of 16-day-old fetal rat stomachs. At this stage of the fetal development the ***gastric*** epithelial cells are undifferentiated. The explants were cultured in DMEM/F-12 medium supplemented with fetal calf serum only, or fetal calf serum combined with either hydrocortisone or pentagastrin. Morphological characterization by means of light microscopy, dye staining and immunostaining was used to identify the growing cells. Both hydrocortisone and pentagastrin accelerated the differentiation towards H,K-ATPase-positive cells, mucus-producing cells and other epithelial cells. H,KATPase-positive cells, which were identified by immunostaining with a monoclonal antibody reacting with the alpha-subunit of the H,K-ATPase, grew on top of the confluent layer of

epithelioid and fibroblastoid cells. With this method in vitro investigations of the mechanisms of proliferation and differentiation of ***gastric*** mucosal cells are possible. Although by different mechanisms, both hydrocortisone and pentagastrin appear to play a regulatory role in these processes.

L2 ANSWER 18 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 12

AN 1997:277254 BIOSIS

DN PREV199799576457

TI Detection of spiral and coccoid forms of Helicobacter ***pylori*** using a murine monoclonal antibody.

AU Cao, J.; Li, Z.-Q.; Borch, K.; ***Mardh, S.***

CS Dep. Cell Biol., Fac. Health Sci., Linkoping Sweden

SO Gastroenterology, (1997) Vol. 112, No. 4 SUPPL., pp. A83.
Meeting Info.: Digestive Disease Week and the 97th Annual Meeting of the American Gastroenterological Association Washington, D.C., USA May 11-14, 1997

ISSN: 0016-5085.

DT Conference; Abstract

LA English

L2 ANSWER 19 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 96:789662 SCISEARCH

GA The Genuine Article (R) Number: VP322

TI SER-27, TYR-10 AND TYR-7 IN THE ALPHA-CHAIN OF PIG STOMACH H+,K+-ATPASE AS CA2+-DEPENDENT PHOSPHORYLATABLE SITES BY INTRINSIC AND EXTRINSIC PROTEIN-KINASES

AU TOGAWA K; KAYA S; SHIMADA A; IMAGAWA T; ***MARDH S***; CORBIN J; KIKKAWA U; TANIGUCHI K (Reprint)

CS HOKKAIDO UNIV, GRAD SCH SCI, SAPPORO, HOKKAIDO 060, JAPAN (Reprint); HOKKAIDO UNIV, GRAD SCH SCI, SAPPORO, HOKKAIDO 060, JAPAN; LINKOPING UNIV, DEPT CELL PHYSIOL, S-58183 LINKOPING, SWEDEN; VANDERBILT UNIV, DEPT MOL PHYSIOL & BIOPHYS, NASHVILLE, TN, 37232; KOBE UNIV, BIOSIGNAL RES CTR, KOBE 657, JAPAN

CYA JAPAN; SWEDEN; USA

SO BIOCHEMICAL AND BIOPHYSICAL RESEARCH COMMUNICATIONS, (23 OCT 1996) Vol. 227, No. 3, pp. 810-815.

ISSN: 0006-291X.

DT Article; Journal

FS LIFE

LA ENGLISH

REC Reference Count: 16

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

When pig stomach membrane H+,K+-ATPase preparations were incubated with [gamma-P-32]ATP, Mg2+ and Ca2+, reversible phosphorylation of specific Tyr and Ser residues in the N-terminal alpha-chain of H+,K+-ATPase occurred without any detectable phosphorylation in other regions of the alpha-chain. Mild tosylphenyla lanyl chloromethyl ketone-trypsin treatment followed by reverse; phase column chromatography yielded three radioactive peptide peaks. The first peak contained both Tyr(10)(P-32) and Tyr(7)(P-32) and the second peak contained Tyr(10)(P-32). The third peak contained Ser(27)(P-32) which was also obtained after trypsin treatment of partially purified H-,K+-ATPase preparations phosphorylated with protein kinase-C + Ca2+ or protein kinase-A. This is the first demonstration of Ca2+-dependent phosphorylation of the alpha-chain of H+,K+-ATPase by

L2 ANSWER 20 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 13

AN 1996:313659 BIOSIS

DN PREV199699036015

TI Interactions between Ca-2+- and cAMP-dependent stimulatory pathways in parietal cells.

AU Li, Zhao-Qi; ***Mardh, Sven (1)***

CS (1) Dep. Cell Biol., Fac. Health Sci., Linkoping Univ., S-581 85 Linkoping Sweden

SO Biochimica et Biophysica Acta, (1996) Vol. 1311, No. 2, pp. 133-142. ISSN: 0006-3002.

DT Article

LA English

AB Isolated rat parietal cells were used to investigate the role of intracellular Ca-2+ in the action of cAMP-dependent secretagogues and cross talk between cAMP- and Ca-2+-dependent stimulatory pathways. Aminopyrine accumulation (an index of acid produced and trapped by the parietal cells), cytosolic free Ca-2+, morphological transformation and cell viability were used to investigate parietal cell function 1,2-bis(2-aminophenoxy)ethane-N,N,N',N'-tetraacetic acid (BAPTA, 10 mu-M). Also, the morphological transformations induced by and stimulation. The increase of cytosolic free Ca-2+ promoted by ***gastrin***, or carbachol, was abolished by the intracellular Ca-2+ chelator dibutytyladenosine 3':5'-cyclic monophosphate (DBcAMP), ***gastrin***, and Sp-adenosine-cyclic-3',5'-monophosphothioate, (Sp-CAMPS) were completely abolished by BAPTA (10 mu-M). In aminopyrine accumulation the action of 1 mM DBcAMP was dose-dependently reduced by BAPTA. The Ca-2+ ionophore A23187 alone, in the range of 1 pM to 1 mu-M, had no effect but it dose-dependently potentiated the action of 1 mM DBcAMP in aminopyrine accumulation. The inhibitory actions of BAPTA on DBcAMP- and histamine-stimulated aminopyrine accumulation were dose-dependently reversed by A23187. Histamine-stimulated protein kinase activity and viability parameters as cellular lactate dehydrogenase (LDH) and trypan blue exclusion were not changed by BAPTA. These results indicated that in isolated parietal cells: (1) the action of cAMP-dependent secretagogues in aminopyrine accumulation and morphological transformation are dependent on cytosolic free Ca-2+; (2) Ca-2+-induced morphological transformation is essential for aminopyrine accumulation; (3) a threshold level of one second messenger is required for stimulation of aminopyrine accumulation by the other second messenger.

L2. ANSWER 21 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 14

AN 1995:154706 BIOSIS

DN PREV199598169006

TI ***Gastrin*** and carbachol require cAMP to elicit aminopyrine accumulation in isolated pig and rat parietal cells.

AU Li, Zhao-Qi; Cabero, Jose Luis; ***Mardh, Sven (1)***

CS (1) Dep. Cell Biol., Fac. Health Sci., Linkoping University, S-581 85 Linkoping Sweden

SO American Journal of Physiology, (1995) Vol. 268, No. 1 PART 1, pp. G82-G89.

ISSN: 0002-9513.

DT Article

LA English

AB The role of endogenous adenosine 3',5'-cyclic monophosphate (cAMP) in the mechanisms of action of ***gastrin*** and carbachol on aminopyrine accumulation in isolated pig and rat parietal cells was investigated. In pig cells, pentagastrin (100 nM) alone stimulated aminopyrine accumulation, an action significantly reduced by the protein kinase A inhibitor Rp-adenosine 3',5'-cyclic monophosphothioate (Rp-cAMP(S); 100 mu-M). In rat cells, ***gastrin*** -17 (100 nM) was incapable of stimulating aminopyrine accumulation, but it potentiated the action of histamine (100 mu-M). Carbachol (10 mu-M) stimulated aminopyrine accumulation and potentiated the action of histamine, and its action was potentiated in a dose-dependent manner by Sp-adenosine 3',5'-cyclic monophosphothioate (SpcAMP(S); a cAMP analogue) in both species. The effect of carbachol was dose dependently reduced by Rp-cAMP(S). The basal cAMP in pig parietal cells was 3.5-fold higher than that in rat parietal cells. Histamine (100 mu-M) and 3-isobutyl-1-methylxanthine (IBMX; 100 mu-M) only slightly elevated the cAMP content (1.2- to 2.9-fold the basal level) in both pig and rat parietal cells. Their combination, however, increased the cAMP level by 8- to 38-fold, but it did not increase aminopyrine accumulation above that elicited by histamine alone. ***Gastrin*** did not alter the cAMP levels in parietal cells of either of the two species. Both ***gastrin*** and carbachol increased cytosolic free Ca-2+ in enriched pig and rat parietal cells. These results indicated that in isolated pig and rat parietal cells 1) secretagogues that elevate intracellular free Ca-2+, such as ***gastrin*** and carbachol, require a certain cAMP level to be effective in stimulating aminopyrine accumulation; 2) a further increase over a certain level of cAMP does not result in an increased aminopyrine accumulation; and 3) parallel activation of Ca-2+- and cAMP-dependent pathways appears to be necessary to effectively elicit aminopyrine accumulation.

L2 ANSWER 22 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 95:80651 SCISEARCH

GA The Genuine Article (R) Number: QB678

TI ***GASTRIN*** AND CARBACHOL REQUIRE CAMP TO ELICIT AMINOPYRINE ACCUMULATION IN ISOLATED PIG AND RAT PARIETAL-CELLS

AU LIZQ; CABEROJL; ***MARDH S (Reprint)***

CS LINKOPING UNIV, FAC HLTH SCI, DEPT CELL BIOL, S-58185 LINKOPING, SWEDEN (Reprint); LINKOPING UNIV, FAC HLTH SCI, DEPT CELL BIOL, S-58185 LINKOPING, SWEDEN

CYA SWEDEN

SO AMERICAN JOURNAL OF PHYSIOLOGY-GASTROINTESTINAL AND LIVER PHYSIOLOGY, (JAN 1995) Vol. 31, No. 1, pp. G82-G89.

ISSN: 0193-1857.

DT Article: Journal

FS LIFE

LA ENGLISH

REC Reference Count: 35

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB The role of endogenous adenosine 3',5'-cyclic monophosphate (cAMP) in the mechanisms of action of ***gastrin*** and carbachol on aminopyrine accumulation in isolated pig and rat parietal cells was investigated. In pig cells, pentagastrin (100 nM) alone stimulated aminopyrine accumulation, an action significantly reduced by the protein kinase A inhibitor Rp-adenosine 3',5'-cyclic monophosphothioate (Rp-cAMP[S]; 100 mu M), In rat cells, ***gastrin*** -17 (100 nM) was incapable of

stimulating aminopyrine accumulation, but it potentiated the action of histamine (100 mu M). Carbachol (10 mu M) stimulated aminopyrine accumulation and potentiated the action of histamine, and its action was potentiated in a dose-dependent manner by Sp-adenosine 3',5'-cyclic monophosphothioate (Sp-cAMP[S]; a cAMP analogue) in both species. The effect of carbachol was dose dependently reduced by Rp-cAMP[S]. The basal cAMP in pig parietal cells was 3.5-fold higher than that in rat parietal cells. Histamine (100 mu M) and 3-isobutyl-1-methylxanthine (IBMX; 100 mu M) only slightly elevated the cAMP content (1.2- to 2.9-fold the basal level) in both pig and rat parietal cells. Their combination, however, increased the cAMP level by 8- to 38-fold, but it did not increase aminopyrine accumulation above that elicited by histamine alone. ***Gastrin*** did not alter the cAMP levels in parietal cells of either of the two species. Both ***gastrin*** and carbachol increased cytosolic free Ca2+ in enriched pig and rat parietal cells. These results indicated that in isolated pig and rat parietal cells 1) secretagogues that elevate intracellular free Ca2+, such as ***gastrin*** and carbachol, require a certain cAMP level to be effective in stimulating aminopyrine accumulation; 2) a further increase over a certain level of cAMP does not result in an increased aminopyrine accumulation; and 3) parallel activation of Ca2+- and cAMP-dependent pathways appears to be necessary to effectively elicit aminopyrine accumulation.

L2 ANSWER 23 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 15 AN 1995:61051 BIOSIS

DN PREV199598075351

TI Positive Correlation between H,K-Adenosine Triphosphatase Autoantibodies and Helicobacter ***pylori*** Antibodies in Patients with Pernicious Anemia.

AU Ma, J.-Y.; Borch, K.; Sjostrand, S. E.; Janzon, L.; ***Mardh, S. (1)***

CS (1) Dep. Cell Biol., Fac. Health Sci., S-581-85 Linkoping Sweden

SO Scandinavian Journal of Gastroenterology, (1994) Vol. 29, No. 11, pp. 961-965.

ISSN: 0036-5521.

DT Article

LA English

AB Background: Helicobacter ***pylori*** is a major cause of ***gastritis***, and the parietal cell H,K-adenosine triphosphatase (ATPase) is a major autoantigen in autoimmune atrophic corpus ***gastritis***, which may eventually lead to pernicious anemia and/or neuropathy. Whether the bacterium induces the autoimmune response is unknown. Methods: By means of enzyme-linked immunosorbent assay the occurrence of antibodies against porcine H,K-ATPase and H. ***pylori*** was determined in sera from 30 patients with pernicious anemia. Results: All sera scored positive against H,K-ATPase, and 25 (83%) scored positive against H. ***pylori*** . The titers of antibodies against both antigen preparations inversely correlated with the duration of disease. A possible common epitope in the antigen preparations was tested with a competition assay. There was no indication of a common epitope in either human or porcine H,K-ATPase and H. ***pylori*** . Conclusions: There was a positive correlation and a high incidence of antibodies against H,K-ATPase and H. ***pylori*** in sera from patients with pernicious anemia. These antibodies recognized different epitopes.

Phis

AN 1994:286849 BIOSIS

DN PREV199497299849

TI Mechanisms of action of secretagogues in isolated pig and rat parietal cells.

AU Li, Z.-Q.; Cabero, J. L.; ***Mardh, S.***

CS Dep. Cell Biol., Fac. Health, Science, Linkoping Sweden

SO Gastroenterology, (1994) Vol. 106, No. 4 SUPPL., pp. A822.

Meeting Info.: 95th Annual Meeting of the American Gastroenterological Association New Orleans, Louisiana, USA May 15-18, 1994

ISSN: 0016-5085.

DT Conference

LA English

L2 ANSWER 25 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 16

AN 1994:496414 BIOSIS

DN PREV199497509414

TI Human ***gastric*** H,K-adenosine triphosphatase beta-subunit is a major autoantigen in atrophic corpus ***gastritis***.

AU Ma, J.-Y.; Borch, K.; ***Mardh, Sven (1)***

CS (1) Dep. Cell Biol., Fac. Health Sci., S-581 85 Linkoping Sweden

SO Scandinavian Journal of Gastroenterology, (1994) Vol. 29, No. 9, pp. 790-794.

ISSN: 0036-5521.

DT Article

LA English

AB Background: Sera from patients with atrophic corpus ***gastritis*** with pernicious anemia frequently contain parietal cell autoantibodies. We have previously demonstrated that the human H,K-adenosine triphosphatase (H,K-ATPase) alpha-subunit constitutes a major autoantigen. The present study investigates whether the human H,K-ATPase beta-subunit is an autoantigen, too. Methods: The gene of the human beta-subunit was expressed in insect cells by a baculovirus expression system. The reactivity of sera from 42 patients towards the recombinant glycoprotein was analyzed by means of an enzyme-linked immunosorbent assay. Results: Thirty-nine of the 42 sera (93%) scored positive. Autoantibody binding in 41 sera (98%) was eliminated when unglycosylated beta-subunit was used as antigen, and antibody binding in the last serum was decreased by 30%. Conclusions: The results indicate that the beta-subunit is indeed a major autoantigen and that carbohydrates are involved in binding of the autoantibodies.

L2 ANSWER 26 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 17

AN 1994:286477 BIOSIS

DN PREV199497299477

TI H,K-ATPase autoantibodies and Helicobacter ***pylori*** antibodies in patients with pernicious anemia.

AU ***Mardh, S.***; Ma, J.-Y.; Janzon, L.; Borch, K.

CS Dep. Cell Biol., Fac. Health Sci., Linkoping Sweden

SO Gastroenterology, (1994) Vol. 106, No. 4 SUPPL., pp. A729.

Meeting Info.: 95th Annual Meeting of the American Gastroenterological.

Association New Orleans, Louisiana, USA May 15-18, 1994

ISSN: 0016-5085.

DT Conference

LA English

L2 ANSWER 27 OF 86 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.DUPLICATE 18

AN 94057929 EMBASE

DN 1994057929

TI Localization of a pernicious anaemia autoantibody epitope on the .alpha.-subunit of human H,K-adenosine triphosphatase.

AU Song Y.-H.; Ma J.-Y.; ***Mardh S.***; Liu T.; Sjostrand S.E.; Rask L.; Borch K.; Huang G.-C.; Barnett P.; McGregor A.M.; Banga J.P.

CS Department Cell Biology, Faculty of Health Sciences, S-581 85 Linkoping, Sweden

SO Scandinavian Journal of Gastroenterology, (1994) 29/2 (122-127). ISSN: 0036-5521 CODEN: SJGRA4

CY Norway

DT Journal; Article

FS 025 Hematology

048 Gastroenterology

LA English

SL English

AB Four cDNA fragments encoding different portions of the .alpha.-subunit of human H,K-adenosine triphosphatase (ATPase) were amplified by means of the polymerase chain reaction technique, ligated into the plasmid pGEX-2T, and expressed as glutathione S-transferase fusion proteins in Escherichia coli. The fragments A (residues 163-313), Ba (residues 360-797), Bb (residues 526-797), and C (residues 822-1031) together encompass 77% of the .alpha.-subunit and cover most of its cytosolic part. Ther reactivities of autoantibodies in the sera from patients with pernicious anaemia with the recombinant fusion proteins were analysed by immunoblotting. One autoantigen epitope was found in the NH2-terminal part of the Ba fragment - that is, between residues 360 and 525. No epitope was detected in the other fragments. The Ba fragment was cleaved off from the glutathione S-transferase fusion protein by the action of thrombin and was then further purified. By means of enzyme-linked immunosorbent assay, 28 of 42 sera (67%) from patients with pernicious anaemia were positive against the purified Ba fragment. The present results provide a final proof that the human H,K-ATPase .alpha.-subunit is a major autoantigen in the parietal cell and that the major epitope is located between residues 360 to 525 on the cytosolic side of the secretory membrane.

L2 ANSWER 28 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 19

AN 1993:184628 BIOSIS

DN PREV199395095078

TI Direct ***gastrin*** action on isolated rat parietal cells induces morphological transformations.

AU Li, Zhao-Qi; Cabero, Jose Luis; Nilsson, B. Ove; ***Mardh, Sven (1)***

CS (1) Uppsala Biomedical Centre, Box 575, S-751 23 Uppsala Sweden

SO Biochimica et Biophysica Acta, (1993) Vol. 1175, No. 3, pp. 250-256. ISSN: 0006-3002.

DT Article

LA English

AB In isolated rat parietal cells, a potentiating effect by ***gastrin***
of the stimulatory action of histamine and dibutyryl-cAMP (DBcAMP) on
aminopyrine accumulation, an index of the acid formed and trapped by the
cells, was recently reported by us (1991, Am. J. Physiol. 261, G621-G627).
In the present study, this mechanism of action of ***gastrin*** was
further investigated. Enriched parietal cells (apprxeq 65% parietal
cells) were incubated under different conditions and processed for

electron microscopy. Morphometric analysis of the micrographs revealed that pentagastrin (100 nM) was as efficient as histamine (100 mu-M) in inducing the formation of vacuolar/canalicular spaces in the parietal cells. In the presence of the histamine H-2-receptor antagonist ranitidine, histamine was ineffective but pentagastrin and ***gastrin*** -17 (G17) maintained their capacity to induce the morphological transformations. By stimulation with pentagastrin plus histamine, the vacuolar/canalicular volume was 2-fold higher than by stimulation separately with each one of the secretagogues. G-17 (100 nM) alone was ineffective but potentiated the maximal (14C)aminopyrine accumulation obtained with 100 mu-M histamine in mucosal cells (apprxeq 25-35% parietal cells). Ranitidine blocked both histamine- and histamine plus G-17-stimulated aminopyrine accumulation. G-17 potentiated also the stimulation by 1 mM dibutyryl-cyclic AMP but this was not inhibited by ranitidine. Pentagastrin (100 nM) increased the basal (14C)glucose oxidation in mucosal cells by 30%. This increase was not blocked by ranitidine which, however, abolished the histamine-stimulated glucose oxidation. Incubation of the cells with pentagastrin plus histamine resulted in a glucose oxidation which equaled the sum of the values. obtained by each one of the agents. These results indicate that ***gastrin***, acting directly on the parietal cells, potentiates the action of histamine on aminopyrine accumulation by increasing the vacuolar/canalicular spaces, a process that is reflected in the metabolic activity of the cells. Thus a major effect of ***gastrin*** at the parietal cell level appears to be the induction of a morphology which is characteristic of stimulated cells rather than a direct activation of ion-transport mechanisms.

L2 ANSWER 29 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 20

AN 1993:386071 BIOSIS

DN PREV199396061371

TI ***Gastrin*** action on aminopyrine accumulation in isolated pig parietal cells requires cAMP.

AU Cabero, Jose Luis; Li, Zhao-Qi; ***Mardh, Sven (1)***

CS (1) Dep. Cell Biol., Fac. Health Sci., S-581 85 Linkoping Sweden

SO Biochimica et Biophysica Acta, (1993) Vol. 1177, No. 3, pp. 245-252. ISSN: 0006-3002.

DT Article

LA English

AB The mechanism of action of ***gastrin*** on pig parietal cells was investigated. The aminopyrine accumulation technique was used to estimate acid production in ***gastric*** mucosal cells, containing 10-20% parietal cells, and in enriched parietal cells, containing 65-95% parietal cells. The ***gastrin*** analogue pentagastrin stimulated aminopyrine accumulation in a dose-dependent fashion irrespective of the proportion of non-parietal cells present. The apparent EC-50 for pentagastrin was 5 nM and the maximally effective concentration was 100 nM. The histamine H-2-receptor antagonist ranitidine did not affect the action of pentagastrin. The stimulatory effects of various doses of histamine on aminopyrine accumulation in highly enriched parietal cells were potentiated by the inclusion of 100 nM pentagastrin in the incubation medium. In another series of experiments using mucosal cells, the action of effective doses of pentagastrin were potentiated by the phosphodiesterase inhibitor isobutylmethyl xanthine (IBMX), which alone elicited an aminopyrine accumulation equal to 50% of that obtained by 100

mu-M histamine. When ranitidine (100 mu-M) was included, the action of IBMX was almost completely abolished. However, the dose-response curve for pentagastrin in the presence of ranitidine plus IBMX was similar to that obtained in the absence of IBMX. Dibutyryl-cAMP (DBcAMP, 1 mM) in the presence of ranitidine (100 mu-M) also potentiated the action of all effective doses of pentagastrin on mucosal cells. The protein kinase A inhibitor Rp-cAMPS, present at 500 mu-M in the incubation medium, significantly reduced the action of each effective concentration of pentagastrin on aminopyrine accumulation in enriched parietal cells. These results in pig parietal cells were interpreted as indicative of: (i) an action of ***gastrin*** exerted directly on the parietal cells; (ii) elevation of intracellular cAMP having a permissive role in the action of ***gastrin*** on aminopyrine accumulation.

L2 ANSWER 30 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1993:310169 BIOSIS

DN PREV199345016694

TI ***Gastric*** hydrogen, potassium-ATPase and mRNA-level in the rat after long term treatment with omeprazole, lansoprazole or pantoprazole.

AU Fryklund, J. (1); Torven, A.; Stalbom, B.-M.; Cabero, J. L.; ***Mardh, ***

*** S.***; Lundberg, L.

CS (1) Astra Hassle AB, Molndal Sweden

SO Gastroenterology, (1993) Vol. 104, No. 4 SUPPL., pp. A82.
Meeting Info.: 94th Annual Meeting of the American Gastroenterological Association Boston, Massachusetts, USA May 15-21, 1993
ISSN: 0016-5085.

DT Conference

LA English

L2 ANSWER 31 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 93:261313 SCISEARCH

GA The Genuine Article (R) Number: KX957

TI ***GASTRIC*** H,K-ATPASE AND MESSENGER RNA-LEVELS IN THE RAT AFTER LONG-TERM TREATMENT WITH OMEPRAZOLE, LANSOPRAZOLE OR PANTOPRAZOLE

AU FRYKLUND J (Reprint); TORVEN A; STALBOM B M; CABERO J L; ***MARDH S***
; LUNDBERG L

CS ASTRA HASSLE AB, MOLNDAL, SWEDEN; UNIV UPPSALA, DEPT MED & PHYSIOL CHEM, S-75105 UPPSALA, SWEDEN

CYA SWEDEN

SO GASTROENTEROLOGY, (APR 1993) Vol. 104, No. 4, Supp. S, pp. A82. ISSN: 0016-5085.

DT Conference; Journal

FS LIFE; CLIN

LA ENGLISH

REC No References

L2 ANSWER 32 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 21

AN 1992:255992 BIOSIS

DN BA93:132317

TI EFFECTS OF ***GASTRIN*** ON CYTOSOLIC FREE CALCIUM IN INDIVIDUAL ACID-SECRETING RAT PARIETAL CELLS.

AU CABERO J L; GRAPENGIESSER E; GYLFE E; LI Z-Q; ***MARDH S***

CS DEP. MED. PHYSIOL. CHEM., BOX 575, UPPSALA UNIV., UPPSALA S-751 23, SWED.

SO BIOCHEM BIOPHYS RES COMMUN, (1992) 183 (3), 1097-1102. CODEN: BBRCA9. ISSN: 0006-291X.

FS BA; OLD

LA English

AB The effects of ***gastrin*** on cytosolic free Ca2+([Ca2+]i) in single, isolated rat ***gastric*** parietal cells were investigated using the fluorescent probe Fura-2 and digital image analysis. [Ca2+]i was increased by ***gastrin*** (100 nM) in apprxeq. 30% of the parietal cells, which were identified by using either the fluorescent probe acridine orange or a parietal cell-specific monoclonal antibody. In the dominant pattern observed, [Ca2+]i was elevated 50-150% and returned within 1-2 min to a value 30-60% over the basal, which was sustained until withdrawal of the stimulant or addition of the ***gastrin*** inhibitor L-365,260 (1 mu.M). The second, but not the first phase, was abolished in the absence of extracellular Ca2+. The results indicate the existence of functional ***gastrin*** receptors in a subpopulation of rat parietal cells.

L2 ANSWER 33 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 22

AN 1992:235380 BIOSIS

DN BA93:123405

TI PRODUCTION OF MONOCLONAL ANTIBODIES AGAINST ***GASTRIC*** PARIETAL CELL ANTIGENS.

AU CABERO J L; SASAKI T; SONG Y-H; HOLMDAHL R; ***MARDH S***

AB Two mice DBA/1 were each immunized with a single injection of one million

CS DEP. MED. AND PHYSIOL. CHEM., BIOMED. CENTRE, UPPSALA UNIV., P.O. BOX 575, S-751 23 UPPSALA, SWEDEN.

SO ACTA PHYSIOL SCAND, (1992) 144 (3), 369-378. CODEN: APSCAX. ISSN: 0001-6772.

FS BA; OLD

LA English

enriched parietal cells in the hind foot pads. Monoclonal antibodies to be used as research tools in studies on regulatory mechanisms in ***gastric*** parietal cells were obtained after fusion of mouse myeloma cells (SP2) with cells from the popliteal lymph nodes of the mice. Twelve hybridomas produced antibodies reactive with structures only present in parietal cells as assessed by immunohistochemistry of oxyntic mucosa sections. Three hybridomas were subcloned and the antibodies produced by them, designated as PC4, PC8, and PC117, were characterized. In an enzyme-linked immunosorbent assay, all antibodies reacted with H,K-ATPase-containing vesicles. The antibody PC8 recognized a 94 kDa protein after immunoblotting of H,K-ATPase-containing vesicles and all antibodies precipitated a 94 kDa protein from [125I]H,K-ATPase-containing vesicles. The antibodies PC4 and PC117 recognized extracellular structures with a polarized distribution in viable, purified parietal cells. The results suggest that the structure recognized by all three antibodies is the .alpha.-subunit of the H,K-ATPase. The antibodies produced by another hybridoma, PC43, recognized a structure present in parietal and surface epithelial cells of the oxyntic mucosa. In an enzyme-linked immunosorbent assay, they reacted with a high-activity carbonic anhydrase which had been affinity-purified from pig oxyntic mucosa and they recognized a 30 kDa protein after immunoblotting. Thus, monoclonal antibodies against both intracellular and extracellular parietal cell structures were obtained after immunization with a small number of parietal cells.

L2 ANSWER 34 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 23 AN 1993:43717 BIOSIS

DN PREV199344020567

TI A continuous flow techniques for analysis of the stoichiometry of the ***gastric*** proton, potassium-ATPase.

AU ***Mardh, Sven (1)***; Norberg, L.

CS (1) Dep. Med. and Physiological Chem., Biomed. Centre, Uppsala Univ., Box 575, S-751 23 Uppsala

SO Acta Physiologica Scandinavica Supplementum, (1992) Vol. 0, No. 607, pp. 259-263.

Meeting Info.: Satellite Symposium of the 15th International Congress of Biochemistry on Ion Pumps, Structure and Mechanism, Gothenburg, Sweden, August 12-14, 1991. ACTA PHYSIOL SCAND SUPPL ISSN: 0302-2994.

DT Article

LA English

L2 ANSWER 35:OF 86 MEDLINE

AN 93080083 MEDLINE

DN 93080083

TI A continuous flow technique for analysis of the stoichiometry of the ***gastric*** H,K-ATPase.

AU ***Mardh S***; Norberg L

CS Department of Medical and Physiological Chemistry, Uppsala University, Sweden..

SO ACTA PHYSIOLOGICA SCANDINAVICA. SUPPLEMENTUM, (1992) 607 259-63. Journal code: 1UF. ISSN: 0302-2994.

CY ENGLAND: United Kingdom

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 199303

AB A continuous flow method was developed for determining the stoichiometry of the ***gastric*** proton pump H,K-ATPase in its hydrolysis of ATP, translocation of H+ and the K+ congener 86Rb+. H.K-ATPase-containing vesicles which had been isolated from pig ***gastric*** mucosa were incubated at 37 degrees C for 2 h in 150 mM 86RbCl, 0.5 mM EGTA and 3 mM Mes-buffer adjusted to pH 6.1 with Tris, and then applied to a 0.45 micron pore size filter. The immobilized vesicles were superfused with 0.15 mM Mes/Tris buffer, pH 6.1, containing 150 mM choline-Cl and 0.2 mM MgCl2. After the medium was changed to one containing 0.1 mM ATP, the amounts and rates of H+ uptake, 86Rb+ efflux, and ATP hydrolysis were measured in fractions collected after the filter. The initial ratio of transported Rb+ to hydrolysed ATP gave values of 0.96 + - 0.26 (mean + - SD, n = 28). The initial ratio of ATP-dependent Rb+ efflux to H+ uptake gave values of 0.92 \pm 0.28 (mean \pm SD, n = 28). The MgATPase activity was measured in vesicles which had been incubated with choline-Cl instead of RbCl. In the initial fractions used for calculation of the stoichiometry, the MgATPase activity was 15.8% +/- 8.7 (mean +/- S.D.) of the maximal ATPase activity obtained with Rb(+)-loaded vesicles. The MgATPase may be an intrinsic activity of the H,K-ATPase. However, whether corrections were made for the MgATPase or not, it had only marginal effects on the calculations of the stoichiometry of the pump. (ABSTRACT TRUNCATED AT 250 WORDS)

L2 ANSWER 36 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 92:603727 SCISEARCH

GA The Genuine Article (R) Number: JT268

TI A CONTINUOUS-FLOW TECHNIQUE FOR ANALYSIS OF THE STOICHIOMETRY OF THE ***GASTRIC*** H.K-ATPASE

AU ***MARDH S (Reprint)***; NORBERG L

CS UNIV UPPSALA, CTR BIOMED, DEPT MED & PHYSIOL CHEM, BOX 575, S-75123 UPPSALA, SWEDEN (Reprint)

CYA SWEDEN

SO ACTA PHYSIOLOGICA SCANDINAVICA, (1992) Vol. 146, Supp. 607, pp. 259-263. ISSN: 0001-6772.

DT Article; Journal

FS LIFE

LA ENGLISH

REC Reference Count: 12

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

A continuous flow method was developed for determining the stoichiometry of the ***gastric*** proton pump H,K-ATPase in its hydrolysis of ATP, translocation of H+ and the K+ congener Rb-86+. H,K-ATPase-containing vesicles which had been isolated from pig ***gastric*** mucosa were incubated at 37-degrees-C for 2 h in 150 mM (RbCl)-Rb-86, 0.5 mM EGTA and 3 mM Mes-buffer adjusted to pH 6.1 with Tris, and then applied to a 0.45 mum pore size filter. The immobilized vesicles were superfused with 0.15 mM Mes/Tris buffer, pH 6.1, containing 150 mM choline-Cl and 0.2 mM MgCl2. After the medium was changed to one containing 0.1 mM ATP, the amounts and rates of H+ uptake, Rb-86+ efflux, and ATP hydrolysis were measured in fractions collected after the filter. The initial ratio of transported Rb+ to hydrolysed ATP gave values of 0.96 +/-0.26 (mean +/-SD, n = 28). The initial ratio of ATP-dependent Rb+ efflux to H+ uptake gave values of 0.92 + - 0.28 (mean +/- SD, n = 28). The MgATPase activity was measured in vesicles which had been incubated with choline-Cl instead of RbCl. In the initial fractions used for calculation of the stoichiometry, the MgATPase activity was 15.8% +/- 8.7 (mean +/- S.D.) of the maximal ATPase activity obtained with Rb+-loaded vesicles. The MgATPase may be an intrinsic activity of the H.K-ATPase. However, whether corrections were made for the MgATPase or not, it had only marginal effects on the calculations of the stoichiometry of the pump. Thus, simultaneous measurements of Rb-86+ efflux, H+ uptake and ATP hydrolysis in immobilized ***gastric*** vesicles gave a stoichiometry of the pump close to a 1:1:1 ratio.

L2 ANSWER 37 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1993:107592 BIOSIS

DN PREV199344049992

TI Effects of ***gastrin*** on isolated rat parietal cells.

AU Li, Zhao-Qi; Cabero, Jose Luis; ***Mardh, Sven***

CS Dep. Med. and Physiological Chemistry, Uppsala Univ., Uppsala Sweden

SO Acta Physiologica Scandinavica Supplementum, (1992) Vol. 0, No. 608, pp. 160.

Meeting Info.: XX Nordic Congress of Physiology and Pharmacology Copenhagen, Denmark August 16-19, 1992 ISSN: 0302-2994.

DT Conference

LA English

L2 ANSWER 38 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 24

AN 1992:233887 BIOSIS

DN BA93:121912

- TI TWO-DIMENSIONAL CRYSTALS OF MEMBRANE-BOUND ***GASTRIC*** PROTON POTASSIUM ATPASE.
- AU HEBERT H; XIAN Y; HACKSELL I; ***MARDH S***
- CS CENT. STRUCTURAL BIOCHEMISTRY, KAROLINSKA INST., NOVUM, S-141 57 HUDDINGE, SWED.
- SO FEBS (FED EUR BIOCHEM SOC) LETT, (1992) 299 (2), 159-162. CODEN: FEBLAL. ISSN: 0014-5793.
- FS BA; OLD
- LA English
- AB Two-dimensional crystallization of membrane-bound H,K-ATPase (EC 3.6.1.36) in vesicle preparations from parietal cells of hog ****gastric*** mucosa was induced by an imidazole buffer containing Mg2+ and VO3- ions. A continuous reorganization of the protein molecules started within a few hours by the formation of linear arrays. At later stages confluent two-dimensional crystals were formed. Electron microscopy and image processing showed that these were of a single tetragonal type. The asymmetric unit consisted of one pear-shaped protein domain corresponding to a H,K-ATPase protomer. Through stain-deficient contact regions four adjacent protein units were connected forming a tetrameric structure.
- L2 ANSWER 39 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 25
- AN 1992:5300 BIOSIS
- DN BA93:5300
- TI OCCURRENCE OF AUTOANTIBODIES AGAINST INTRINSIC FACTOR PROTON POTASSIUM ATPASE AND PEPSINOGEN IN ATROPHIC ***GASTRITIS*** AND RHEUMATOID ARTHRITIS.
- AU ***MARDH S***; MA J Y; SONG Y H; ALY A; HENRIKSSON K
- CS DEP. MED. PHYSIOL. CHEM., BIOMED. CENTER, UPPSALA UNIV., BOX 575, S-751 23 UPPSALA, SWED.
- SO | SCAND J GASTROENTEROL, (1991) 26 (10), 1089-1096. CODEN: SJGRA4. ISSN: 0036-5521.
- FS BA; OLD
- LA English
- AB The occurrence of autoantibodies against intrinsic factor, H,K-ATPase, and pensinogen was analysed by means of enzyme-linked immunosorbent assay in three groups of sera. Group 1 comprised sera from 14 rheumatoid arthritis patients with normal acid secretion; group 2, sera from 18 rheumatoid arthritis patients with reduced acid secretion; and group 3, sera from 11 patients with pernicious anaemia or achylia. Groups 1 and 2 were rheumatoid factor-positive, and group 3 was negative. Intrinsic factor autoantibodies were low in groups 1 and 2. In group 3, 9 of the 11 sera (82%) scored positive. The highest titres of H,K-ATPase and pepsinogen autoantibodies were found in groups 2 and 3. Only one serum in group 1 scored positive against H,K-ATPase, and two against pepsinogen, whereas corresponding values were 11 (61%) and 7 (39%) in group 2, and 10 (91%) and 6 (55%) in group 3. Autoantibodies against H,K-ATPase from a pool of patient sera recognized both the .alpha.- and .beta.-subunits of the enzyme. The present results support the hypothesis of an autoimmune disease overlap between non-organ-specific rheumatoid arthritis and organ-specific pernicious anaemia.
- L2 ANSWER 40 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 26
- AN 1992:25653 BIOSIS
- DN BA93:14928
- TI ***GASTRIN*** POTENTIATES HISTAMINE-STIMULATED AMINOPYRINE

ACCUMULATION IN ISOLATED RAT PARIETAL CELLS.

AU CABERO JL; LI Z-Q; ***MARDH S***

CS BIOMEDICAL CENTRE, UPPSALA UNIVERSITY, BOX 575, S-751 23 UPPSALA, SWED.

SO AM J PHYSIOL, (1991) 261 (4 PART 1), G621-G627. CODEN: AJPHAP. ISSN: 0002-9513.

FS BA; OLD

LA English

AB Rat ***gastric*** mucosal cells, containing 25-35% parietal cells, were obtained by a modified isolation procedure involving protease, ethylene glycol-bis(.beta.-aminoethyl ether)-N,N,N',N'-tetraacetic acid, and mechanical treatments. Parietal cell responsiveness to secretagogues was assessed by the accumulation of the weak base [14C]aminopyrine in intracellular acidic compartments. Histamine, without phosphodiesterase inhibitors, dose dependently stimulated aminopyrine accumulation with an effective concentration producing 50% of maximal response of 13 .mu.M and a maximal effective dose of 100 .mu.M. Pentagastrin and rat ***gastrin*** -17 alone were ineffective but potentiated dose dependently the action of 100 .mu.M histamine. The mean potentiating effect varied from 32 to 70% for 100 nM pentagastrin and from 36 to 95% for 100 nM rat ***gastrin*** -17. Pentagastrin (100 nM) also potentiated the effect of 1 mM dibutyryrladenosine 3',5'-cyclic monophosphate (cAMP) by 44%, but it did not increase further the stimulation by carbachol. The potentiating effect of pentagastrin on histamine- and dibutyryl cAMP-stimulated aminopyrine accumulation was also observed after enrichment of parietal cells to 65-85%. The endogenous histamine was insufficient to stimulate acid production. Therefore ***gastrin*** appears to have a direct

L2 ANSWER 41 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 27

AN 1991:362541 BIOSIS

action also in rat parietal cells.

DN BA92:50766

TI PEPTIC ULCER DISEASE ABSENCE OF ANTIBODIES STIMULATING THE HISTAMINE SENSITIVE ADENYLATE CYCLASE OF ***GASTRIC*** MUCOSAL CELLS.

AU BURMAN P; ***MARDH S***; LOOF L; NAESDAL J; KARLSSON F A

CS DEP. INTERNAL MEDICINE, UNIVERSITY HOSPITAL, S-751 85 UPPSALA, SWED.

SO GUT, (1991) 32 (6), 620-623.

CODEN: GUTTAK. ISSN: 0017-5749.

FS BA: OLD

LA English

AB The possible presence of parietal cell stimulating antibodies was examined in sera from 57 patients with relapsing ulcer disease. The sera were obtained at the time of symptomatic relapse and all patients had ulcers confirmed by endoscopy. A sensitive assay based on adenosine 3':5' cyclic monophosphte (cAMP) production in isolated procine ***gastric*** mucosal cells was used as a measure. cAMP production increased up to four hours of incubation and was histamine responsive; an approximately 20-fold increase was found with histmaine 10-4 mol/l. Sera from both patients and healthy control subjects showed some inhibitory effect on basal cAMP production compared with incubation in medium only, whereas immunoglobulin preparations had a weaker non-specific effect. No stimulation was found when the patients' sera and immunoglobulins (up to a concnetration of 6 mg/ml) were examined. These results that ***gastric*** acid hypersecretion in duodenal ulcer disease is not an effect of histamine receptor stimulating antibodies. The data thus argue against a recent hypothesis that severe chronic ulcer disease in some patients has an

L2 ANSWER 42 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 28

AN 1992:468999 BIOSIS

DN BR43:90349

TI EFFECTS OF ***GASTRIN*** ON ISOLATED PARIETAL CELLS.

AU ***MARDH S***; CABERO J L; LI Z-Q

CS DEP. MED. PHYSIOLOGY CHEM., UPPSALA UNIV. BIOMEDICAL CENTRE, BOX 575, S-751 23 UPPSALA, SWED.

SO HAKANSON, R. AND F. SUNDLER (ED.). FERNSTROM FOUNDATION SERIES, VOL. 15. THE STOMACH AS AN ENDOCRINE ORGAN; 18TH ERIC K. FERNSTROM SYMPOSIUM, LUND, SWEDEN, MAY 21-23, 1990. XIX+548P. ELSEVIER SCIENCE PUBLISHERS B.V.:

AMSTERDAM, NETHERLANDS; (DIST. IN THE USA AND CANADA BY ELSEVIER SCIENCE PUBLISHING CO., INC.: NEW YORK, NEW YORK, USA). ILLUS. (1991) 0 (0), 253-266.

CODEN: FFOSDF. ISSN: 0167-7004. ISBN: 0-444-81377-2.

DT Conference

FS BR; OLD

LA English

L2 ANSWER 43 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 29

AN 1992:4790 BIOSIS

DN BA93:4790

TI COMPLEMENTARY DNA CLONING OF THE BETA-SUBUNIT OF THE HUMAN ***GASTRIC*** PROTON POTASSIUM ATPASE.

AU MA J-Y; SONG Y-H; SJOSTRAND S E; RASK L; ***MARDH S***

CS DEP. MED. PHYSIOLOGICAL CHEM., BIOMED. CENTRE, BOX 575, S-751 23 UPPSALA, SWED.

SO BIOCHEM BIOPHYS RES COMMUN, (1991) 180 (1), 39-45. CODEN: BBRCA9. ISSN: 0006-291X.

FS BA; OLD

LA English

AB A full-length cDNA clone encoding the human ***gastric*** H,K-ATPase (EC 3. 6. 1. 36) .beta.-subunit was isolated from a human ***gastric*** mucosal lambda.gt10 library using oligonucleotide probes which were based on the cDNA sequence from rat and rabbit H,K-ATPase .beta.-subunits. The insert was 1407 bp in length and encoded a polypeptide of 291 amino acids with a MW = 33,367 Da. It exhibited 84.2%, 85.6% and 81.3% identity to the H,K-ATPase .beta.-subunits of rabbit, pig and rat, respectively.

L2 ANSWER 44 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 91:511129 SCISEARCH

GA The Genuine Article (R) Number: GD854

TI EFFECTS OF ***GASTRIN*** ON ISOLATED RAT PARIETAL-CELLS

AU CABERO J (Reprint); LIZQ; BANDYOPADHYAYS; ***MARDH S***

CS UNIV UPPSALA, CTR BIOMED, DEPT MED & PHYSIOL CHEM, S-75123 UPPSALA, SWEDEN CYA SWEDEN

SO ACTA PHYSIOLOGICA SCANDINAVICA, (1991) Vol. 143, No. 1, pp. A15.

DT Conference; Journal

FS LIFE

LA ENGLISH

REC Reference Count: 3

L2 ANSWER 45 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 30 AN 1991:112070 BIOSIS

DN BA91:59460

TI A CONTINUOUS-FLOW TECHNIQUE FOR ANALYSIS OF STOICHIOMETRY AND TRANSPORT KINETICS OF ***GASTRIC*** HYDROGEN POTASSIUM ATPASE.

AU NORBERG L; ***MARDH S***

CS DEP MED. PHYSIOLOGICAL CHEM., BIOMEDICAL CENTRE, UPPSALA UNIV., BOX 575, S-751 23 UPPSALA, SWED.

SO ACTA PHYSIOL SCAND, (1990) 140 (4), 567-574. CODEN: APSCAX. ISSN: 0001-6772.

FS BA; OLD

LA English

AB A continuous-flow method was developed for determining the stoichiometry of the ***gastric*** proton pump H,K-ATPase (EC 3.6.1.36) in its hydrolysis of ATP and translocation of H+ and the K+ congener 86Rb+. H,K-ATPase-containing vesicles which had been isolated from pig ***gastric*** mucosa were incubated at 37.degree. C for 2 h in 150 mM 86RbCl, 0.5 mM ethylenebis(oxyethylenenitrilo)tetra-acetic acid and 3 mM 2-(N-morpholino)ethane sulphonic acid (Mes) adjusted to pH 6.1 with Tris, and then applied onto a 0.45 .mu.m pore size cellulose acetate filter. The immobilized vesicles were superfused with 0.15 mM Mes/Tris buffer, pH 6.1, containing 150 mM choline chloride and 0.2 mM MgCl/2. After changing to a medium containing 0.1 mmM ATP, the amounts and rates of H+ uptake, 86Rb+ efflux and ATP hydrolysis were measured. The initial ratio of Rb+ transported to ATP hydrolysed gave values of 0.96 .+-. 0.26 (mean .+-. SD, n = 28). The initial ratio of ATP-dependent Rb+ efflux to H+ uptake gave values of 0.92 .+- 0.28 (mean .+- SD, n = 28). The Mg-ATPase activity was measured in vesicles which had been incubated with choline chloride instead of RbCl. This activity was 15.8 .+-. 8.7% (mean .+-. SD) of the total ATPase activity in the initial fractions used for calculation of the stoichiometry. It is argued that this Mg-ATPase may be an intrinsic activity of the H,K-ATPase and that the relation between these activities is dependent on the amount of K+ (or Rb+) present in the assay. However, whether corrections were made for this Mg-ATPase or not, it had only marginal effects on the calculations of the stoichiometry of the pump. Thus simultaneous measurements of 86Rb+ efflux, H+ uptake and ATP hydrolysis in immobilized ***gastric*** vesicles gave a stoichiometry of the pump close to a 1:1:1 ratio. These results indicate that the pump is non-electrogenic.

L2 ANSWER 46 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 31

AN 1991:47194 BIOSIS

DN BA91:25475

ȚI CALCIUM AND CALMODULIN STIMULATE PHOSPHOLIPASE A-2 AND FUSION OF HYDROGEN POTASSIUM ATPASE-CONTAINING MEMBRANE VESICLES ISOLATED FROM PIG ****GASTRIC**** MUCOSA.

AU OLAISSON H; ***MARDH S***; ARVIDSON G

CS DEP. MED. PHYSIOL. CHEM., BIOMED. CENT., UNIV. UPPSALA, BOX 575, S-751 23 UPPSALA, SWEDEN.

SO ACTA PHYSIOL SCAND, (1990) 140 (3), 393-400. CODEN: APSCAX. ISSN: 0001-6772.

FS BA; OLD

LA English

AB Fusion of pig ***gastric*** H,K-ATPase- and phospholipase
A2-containing vesicles in vitro was studied by electron microscopy and by
monitoring the change in fluorescence of octadecyl rhodamine B-labelled
vesicles. Ca2+ stimulated fusion of vesicles, and the fusion rate showed a

positive correlation with the activity of the phospholipase A2. Both the Ca2+-stimulated fusion rate and the Ca2+-dependent phospholipase A2 activity were futher enhanced by the presence of calmodulin. The present results supported our previous findings (Olaisson et al. 1990) and further indicate that the phospholipase A2 associated with the H,K-ATPase-containing membranes might play a central role in membrane fusion processes in the stimulated parietal cell.

L2 ANSWER 47 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 32

AN 1991:52202 BIOSIS

DN BA91:30483

TI OCCURRENCE OF PHOSPHOLIPASE A-2 AND LYSOPHOSPHOLIPASE IN A ***GASTRIC***
HYDROGEN POTASSIUM ATPASE-CONTAINING MEMBRANE FRACTION AND THE FORMATION
OF LYSOPHOSPHATIDYLCHOLINE IN STIMULATED PIG PARIETAL CELLS.

AU OLAISSON H; ARVIDSON G; MA J-Y; ***MARDH S***

CS DEP. MED. PHYSIOL. CHEM., BIOMED. CENT., UPPSALA UNIV., BOX 575, S-751 23 UPPSALA, SWEDEN.

SO ACTA PHYSIOL SCAND, (1990) 140 (3), 383-392. CODEN: APSCAX. ISSN: 0001-6772.

FS BA; OLD

LA English

AB A membrane fraction containing H,K-ATPase (EC 3.6.1.36) was prepared from pig ***gastric*** mucosa and found to contain phospholipase A2 (EC 3.1.1.4) and lysophospholipase (EC 3.1.1.5) activities. Washing the membranes decreased their protein content by 25%. Recovery profiles of H,K-ATPase, phospholipase A2 and lysophospholipase were similar for membranes washed either with water or with 0.15 or 1.5M KCl. Nearly identical distribution profiles were obtained for the three enzyme activities after centrifugation of washed vesicle membranes on a linear sucrose gradient. The phospholipase A2 activity was stimulated by calcium and increased further in the presence of calmodulin. The amount of cellular radioactively labelled lysophosphatidylcholine was doubled upon cholinergic stimulation of isolated parietal cells prelabelled with [3H]glycerol or 32P. The liberated lyso[32P]phosphatidylcholine had its acyl chain in the sn-1 position, which implies an activation of a phospholipase A2. These findings indicate that secretagogues which increase the cytosolic Ca2+ concentration, i.e. acetylcholine, histamine and ***gastrin***, may activate a phospholipase A2 in the parietal cell.

L2 ANSWER 48 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 33

AN 1990:216332 BIOSIS

DN BA89:113622

TI BINDING OF CHOLECYSTOKININ AND SOMATOSTATIN TO ISOLATED PORCINE ***GASTRIC*** MUCOSAL CELLS AND EFFECTS ON AMINOPYRINE UPTAKE.

AU SJODIN L; ENGLUND L J; ***MARDH S***

CS SOCIALSTYRELSENS LAKEMEDELSAVDELNING, PO BOX 607, S-751 25 UPPSALA, SWED.

SO ACTA PHYSIOL SCAND, (1990) 138 (3), 369-376. CODEN: APSCAX. ISSN: 0001-6772.

FS BA; OLD

LA English

AB Mucosal cells were prepared by enzymatic digestion of procine

gastric mucosa with pronase and collagenase. The resulting cell
suspension contained 10-15% parietal cells, which responded to histamine
stimulation by an up to 20-fold increase in [14C]aminopyrine accumulation

over control levels. Cholecystokinin-8 (CCK-8) evoked a more moderate stimulation of [14C]aminopyrine accumulation, whereas somatostatin inhibited histamine-stimulated accumulation. Parietal cells were enriched by elutriation and isopycnic centrifugation on density gradients of Percoll. A fraction with 60% parietal cells bound approximately three times more iodinated CCK-8 than a fraction containing 70% non-parietal cells. Binding of [1251]BH-CCK-8 to preparations containing 30-60% parietal cells was specifically inhibited to about 50% by 10-9 M unlabelled CCK-8 but not by bombesin. Cell fractions containing about 30% parietal cells also bound [1251]somatostatin. Unlabelled somatostatin at 10-9 M inhibited tracer binding by about 50%, while CCK-8 did not affect somatostatin binding to such a preparation. The results suggest the existence of specific receptors for CCK and somatostatin on porcine parietal cells exerting a regulatory influence on acid secretion.

L2 ANSWER 49 OF 86 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 90233623 EMBASE

DN 1990233623

TI The occurrence of auto-antibodies in patients with gastro-duodenal lesions.

AU ***Mardh S.***; Song Y.-H.

CS Department of Medical and Physiological Chemistry, Biomedical Centre, Uppsala-University, Box-575, S-751-23 Uppsala, Sweden

SO Journal of Internal Medicine, Supplement, (1990) 228/732 (77-82). ISSN: 0955-7873 CODEN: JIMSE3

CY United Kingdom

DT Journal; Conference Article

FS 025 Hematology

026 Immunology, Serology and Transplantation

048 Gastroenterology

LA English

SL English

AB The occurrence of auto-antibodies in patients with the autoimmune disease perficious anaemia and in patients with active duodenal ulcers was investigated. In order to characterize antigenic structures, various cellular and subcellular fractions were prepared from pig ***gastric*** mucosa and from a homogenate of duodenal mucosa, By means of an enzyme-linked immunosorbent assay and immunoblotting, both the H+,K+-ATPase and pepsinogen/pepsin were shown to constitute the major antigens. All of the seven pernicious-anaemia sera that were tested contained auto-antibodies against both antigens, and the epitopes of the H+,K+-ATPase were shown to be localized on its cytoplasmic face. In 75% (18-24) of the sera from patients with duodenal ulcers, auto-antibodies were detected when using purified antigens. Six sera reacted with H+,K+-ATPase and twelve reacted with pepsinogen, one reacted with both antigens, and four sera reacted with the duodenal mucosal antigen. The occurrence of auto-antibodies indicates that there is a mucosal lesion and that immunological factors may be involved in the pathogenesis of the disease in some patients.

L2 ANSWER 50 OF 86 MEDLINE

AN 90343918 MEDLINE

DN 90343918

TI The occurrence of auto-antibodies in patients with gastro-duodenal lesions.

AU ***Mardh S***; Song Y H

CS Department of Medical and Physiological Chemistry, Biomedical Centre, Uppsala University, Sweden.

SO JOURNAL OF INTERNAL MEDICINE. SUPPLEMENT, (1990) 732 77-82. Journal code: ABK. ISSN: 0955-7873.

CY ENGLAND: United Kingdom

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 199011

AB The occurrence of auto-antibodies in patients with the autoimmune disease pernicious anaemia and in patients with active duodenal ulcers was investigated. In order to characterize antigenic structures, various cellular and subcellular fractions were prepared from pig ***gastric*** mucosa and from a homogenate of duodenal mucosa. By means of an enzyme-linked immunosorbent assay and immunoblotting, both the H+,K(+)-ATPase and pepsinogen/pepsin were shown to constitute the major antigens. All of the seven pernicious-anaemia sera that were tested contained auto-antibodies against both antigens, and the epitopes of the H+,K(+)-ATPase were shown to be localized on its cytoplasmic face. In 75% (18/24) of the sera from patients with duodenal ulcers, auto-antibodies were detected when using purified antigens. Six sera reacted with H+,K(+)-ATPase and twelve reacted with pepsinogen, one reacted with both antigens, and four sera reacted with the duodenal mucosal antigen. The occurrence of auto-antibodies indicates that there is a mucosal lesion and that immunological factors may be involved in the pathogenesis of the disease in some patients.

L2 ANSWER 51 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 34

AN 1989:313235 BIOSIS

DN BA88:26965

TI PARIETAL CELL ANTIBODIES IN PERNICIOUS ANEMIA INHIBIT PROTON POTASSIUM ATPASE THE PROTON PUMP OF THE STOMACH.

AU BURMAN P; ***MARDH S***; NORBERG L; KARLSSON F A

CS DEP. INTERN. MED., UNIV. HOSP., S-751 85 UPPSALA, SWEDEN.

SO GASTROENTEROLOGY, (1989) 96 (6), 1434-1438. CODEN: GASTAB. ISSN: 0016-5085.

FS BA; OLD

LA English

AB Antibodies to a membrane-bound antigen, localized to the canalicular structures of the parietal cell, are found in most sera of patients with chronic atrophic ***gastritis*** and pernicious anemia. In the present study immunoglobulins containing parietal cell antibodies were found to inhibit the activity of H+,K+-adenosine triphosphatase (EC 3.6.1.36) in a tubulovesicular membrane preparation from porcine ***gastric*** mucosa. The degree of inhibition correlated to the titer of parietal cell antibodies as assessed by an enzyme-linked immunosorbent assay. The specificity of the enzymatic inhibition was confirmed by the lack of effect of parietal cell antibodies on membrane-bound esterase. A possible interaction of parietal cell antibodies with ***gastrin*** binding at the receptor level was investigated in a radioreceptor assay employing 125I- ***gastrin*** 1 and a ***gastric*** mucosal cell suspension from the guinea pig. No blocking capacity was found with immunoglobulins from patients with pernicious anemia as compared with immunoglobulins from healthy controls. The results thus demonstrate a direct inhibitory effect

of parietal cell antibodies on the acid producing H+,K+-adenosine triphosphatase of the parietal cell, but also a lack of interaction with the ***gastrin*** receptor, and indicate that in the development of hypo/achylia H+,K+-adenosine triphosphatase autoantibodies could have a major pathogenic role.

L2 ANSWER 52 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 35

AN 1990:107633 BIOSIS

DN BA89:57124

TI THE EFFECT OF ARACHIDONIC ACID AND ITS METABOLITES ON ACID PRODUCTION IN ISOLATED HUMAN PARIETAL CELLS.

AU JARAMILLO E; ***MARDH S***; GREEN K; PERSSON B; RUBIO C; ALY A

CS SECT. GASTROENTEROL., DEP. MED., KAROLINSKA HOSP., S-104 01 STOCKHOLM, SWED.

SO SCAND J GASTROENTEROL, (1989) 24 (10), 1231-1237. CODEN: SJGRA4. ISSN: 0036-5521.

FS BA; OLD

LA English

AB The effect of arachidonic acid and its metabolites on the histamine-stimulated acid production in human isolated parietal cells provenient from endoscopic biopsies was examined. 14C-aminopyrine (14C-AP) accumulation in the parietal cells was used for evaluation of acid production. Histamine dose-dependently increased AP uptake. Histamine stimulation (taken as 100% at 10-5 M) was significantly inhibited by prostaglandin (PG) E2 to 66 .+-. 7% at 10-8 M, 42 .+-. 8% at 10-6 M, and 13 .+-. 10% at 10-4 M (mean .+-. SEM, n = 10). PGF2.alpha., PGD2, and PGI2 showed significant inhibitory effects only at very high concentrations (10-5-10-4 M). Leukotriene (LT) B4 and LTC4 were without effect. The basal acid production (taken as 0%) was lowered significantly by 10-6 M arachidonic acid to -20 .+-. 7.4% (p < 0.02, n = 10), and the histamine-stimulated (10-6 M) acid production from 100% to 64 .+-. 7.2% (p < 0.001, n = 10). Aspirin (10-3 M) increased basal (45 .+-. 9.6%, p < 0.001, n = 10) and histamine-stimulated (10-6 M) acid production (164 .+-. 16.3%, p < 0.001). It is concluded that PGE2, the major product from arachidonic acid metabolism in the human ***gastric*** mucosa, is a significant inhibitor of the histamine-stimulated human parietal cell and may, in humans, play a role as a local physiologic inhibitor of acid secretion.

L2 ANSWER 53 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 36

AN 1989:493408 BIOSIS

DN BA88:119945

TI CHARACTERIZATION OF ANTIGENIC STRUCTURES IN AUTOIMMUNE ATROPHIC

GASTRITIS WITH PERNICIOUS ANEMIA THE PARIETAL CELL PROTON

POTASSIUM ATPASE AND THE CHIEF CELL PEPSINOGEN ARE THE TWO MAJOR ANTIGENS.

AU ***MARDH S***; SONG Y-H

CS DEP. MED. AND PHYSIOL. CHEM., BIOMEDICAL CENT., UPPSALA UNIV., BOX 575, S-751 23 UPPSALA, SWEDEN.

SO ACTA PHYSIOL SCAND, (1989) 136 (4), 581-588. CODEN: APSCAX. ISSN: 0001-6772.

FS BA; OLD

LA English

AB Using isolated cells and subcellular fractions from pig ***gastric***
mucosa, antigenic structures with specific binding of IgG from sera of
patients with auto-immune atrophic ***gastritis*** were characterized

by means of immunoblotting and enzyme-linked immunosorbent assay. In immunoblotting experiments using mucosal cells as the antigen source, two dominating bands of 94 and 41 kDa were found. The two major antigens were identified as the H,K-ATPase (94 kDa), which constitutes the parietal cell acid pump, and pepsinogen (41 kDa) located in the chief cells. There was also a small but significant binding of antibodies to a preparation of Na,K-ATPase, an enzyme which is about 60% homologous to H,K-ATPase. Commercial preparations of hog ***gastric*** pepsinogen and pepsin bound pernicious anaemia IgG with equal efficacy. When sera from seven patients with the diagnosis pernicious anaemia were tested, all were found to contain auto-against H,K-ATPase as well as pepsinogen. In intact, isolated H,K-ATPase-containing vesicles the cytosolic part of the ATPase molecule is facing the outside of the vesicles. Both intact and trypsinized vesicles were incubated with patient sera and with a monoclonal antibody against H,K-ATPase. Pernicious anaemia IgG was found to bind to a cytosolic, trypsin-resistant structure, but the binding of the monoclonal antibody was lost upon trypsinization. The present results indicate that intracellular structures of the ***gastric*** mucosa, due to cell damage, may be exposed to immune-competent cells, which do not recognize these structures as 'self'.

L2 ANSWER 54 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 37

AN 1990:107457 BIOSIS

DN BA89:56948

TI THE OCCURRENCE OF ***GASTRIC*** AND DUODENAL AUTO-ANTIBODIES IN PEPTIC ULCER DISEASE.

AU SONG Y-H; ***MARDH S***

CS DEP. MED. AND PHYSIOL. CHEM., BIOMEDICAL CENT., UPPSALA UNIV., BOX 575, S-751 23 UPPSALA, SWEDEN.

SO ACTA PHYSIOL SCAND (1989) 137 (4), 535-540. CODEN: APSCAX. ISSN: 0001-6772.

FS BA; OLD

LA English

AB The possible relationship between peptic ulcer and the occurrence of auto-antibodies was investigated by means of an enzyme-linked immunosorbent assay (ELISA). Sera from 24 patients with active duodenal ulcer were analysed using cells and subcellular fractions from pig ***gastric*** and dudodenal mucosa for bindingof immunoglobulins. Four sera (17%) reacted with a homogenate from duodenal mucosa. Nine sera (38%) were found to contain auto-antibodies against ***gastric*** mucosal cells. The cell-reactive auto-antibodies were shown to bind preferentially to parietal cells and chief cells. In these cells the antigens were identified as H, K-ATPase and pepsinogen respectively. Six sera were positive against purified H,K-ATPase; 12 sera were positive against pepsinogen, and only one of these sera reacted with both H,K-ATPase and pepsinogen. The results show that auto-antibodies ar formed in a large fraction of patients (18/24, 75%) with peptic ulcer disease. The present study further demonstrates that enrichment of antigenic structures is required for obtaining a satisfactory sensitivity in the assay.

L2 ANSWER 55 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 38

AN 1989:446052 BIOSIS

DN BA88:94324

TI THE EFFECTS OF VARIOUS ***GASTRINS*** ON INTRACELLULAR FREE CALCIUM IN ISOLATED PIG PARIETAL CELLS.

AU CABERO JL, REHFELD JF, ***MARDH S***

CS DEP. MED. PHYSIOL. CHEM., UPPSALA UNIV., BOX 575, S-751 23 UPPSALA, SWEDEN.

SO ACTA PHYSIOL SCAND, (1989) 136 (3), 301-308. CODEN: APSCAX. ISSN: 0001-6772.

FS BA; OLD

LA English

AB ***Gastrin*** 17 (G17) is a potent stimulant of ***gastric*** acid secretion in vivo. In this study, the effects of G17 and some related on intracellular free Ca2+ in isolated pig parietal cells were studied. Both G17 and the synthetic peptide pentagastrin increased intracellular free Ca2+ in a dose-dependent manner over the concentration range 10-6 to 10-6 M, suggesting a specific action. The EC50 values were 3 .times. 10-8 M for G17 and 8 .times. 10-8 M for pentagastrin. The N-terminal tridecapeptide of G17 [(1-13)G17] did not have any effect on intracellular free Ca2+, nor was it able to inhibit the action of G17. A glycine-extended ***gastrin*** [(5-17)G17-Gly)] elicited a small but significant increase in intracellular free Ca2+ although only at 10-6 M. This increase was approximately 20% of that obtained with a similar concentration of G17. Sequential incubations with (5-17)G17 and G17 showed that both peptides increased the intracellular free Ca2+ through the same mechanisms.

L2 ANSWER 56 OF 86 MEDLINE

AN 89072509 MEDLINE

DN 89072509

TI Effects of some antisecretory drugs on acid production, intracellular free Ca2+, and cyclic AMP production in isolated pig parietal cells.

AU ***Mardh S*** , Song Y H, Wallmark B

CS Dept. of Medical and Physiological Chemistry, Uppsala University, Sweden..

SO SCANDINAVIAN JOURNAL OF GASTROENTEROLOGY, (1988 Oct) 23 (8) 977-82. Journal code: UCS. ISSN: 0036-5521.

CY Norway

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 198903

AB The effects of some inhibitors of acid secretion were tested on isolated, purified pig parietal cells. The cells were stimulated with 10(-4) M histamine, 10(-5) M carbachol, or 10(-7) M pentagastrin. The H,K-ATPase inhibitors SCH 28080 and omeprazole inhibited both the basal and secretagogue-stimulated acid production, as measured by aminopyrine accumulation, irrespective of the type of stimulator used. The IC50 value was 3-5 x 10(-9) M for SCH 28080 and 1-3 x 10(-8) M for omeprazole. Ranitidine inhibited the histamine-stimulated but not the basal acid production. The IC50 value was 2 x 10(-5) M. Stimulation of acid production with carbachol was blocked by pirenzepine, with an IC50 of 6 x 10(-7) M. Pirenzepine (10(-5) M) specifically blocked the carbachol-stimulated increase in cytosolic free Ca2+ in fura-2-loaded cells but not the increase in cytosolic free Ca2+ induced by histamine or pentagastrin. Ranitidine (10(-4) M) prevented the histamine-induced increase in Ca2+ and was also the only one of the four inhibitors which prevented the histamine-stimulated cAMP formation. SCH 28080 (10(-5) M) significantly potentiated the histamine-stimulated increase in cytosolic free Ca2+ and the formation of cAMP, whereas omeprazole (10(-5) M) was without effect.

L2 ANSWER 57 OF 86 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.DUPLICATE 39

AN 89029721 EMBASE

DN 1989029721

TI Inhibition of H,K-ATPase and Na,K-ATPase by DIDS, a disulphonic stilbene derivative.

AU Vega F.V.; Cabero J.L.; ***Mardh S.***

CS Department of Medical and Physiological Chemistry, Biomedical Centre, Uppsala University, S-751 23 Uppsala, Sweden

SO Acta Physiologica Scandinavica, (1988) 134/4 (543-547). ISSN: 0001-6772 CODEN: APSCAX

CY Sweden

DT Journal

FS 002 Physiology

048 Gastroenterology

LA English

SL English

AB Disulphonic stilbenes are effective inhibitors of an anion exchanger which is present in the plasma membranes of many cells (Cabantchik et al. 1978). In the present study, the effects of 4,4'-diisothiocyano-2,2'-disulphonic stilbene acid (DIDS) on the transport activity of the hydrochloric acid pump isolated from pig stomach (H,K-ATPase, EC 3.6.1.36) were tested. Half-maximal inhibition of proton transport carried out by the H,K-ATPase in the isolated vesicles was observed at micromolar concentrations of DIDS. The effects of DIDS on the adenosine-triphosphatase and p-nitrophenylphosphatase activities of isolated H,K-ATPase were also studied and compared with those of the kinetically and structurally related Na,K-ATPase (EC 3.6.1.37). Half-maximal inhibition of the enzymatic activities of both enzymes were observed in the micromolar range of DIDS. The lipid bilayer of the ***gastric*** vesicle membrane is highly asymmetric and the original cytosolic side is facing the outside of the vesicle. Since DIDS does not readily cross the membrane, it is most likely that DIDS exerts its inhibitory effects by modifying the transport ATPases on their cytosolic sides.

L2 ANSWER 58 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 40

AN 1988:178288 BIOSIS

DN BA85:90390

TI MAJOR PARIETAL CELL ANTIGEN IN AUTOIMMUNE ***GASTRITIS*** WITH PERNICIOUS ANEMIA IS THE ACID-PRODUCING PROTON POTASSIUM ATPASE OF THE STOMACH.

AU KARLSSON F A; BURMAN P; LOOF L; ***MARDH S***

CS DEP. INTERN. MED., UNIV. HOSP., S-751 85 UPPSALA, SWED.

SO J CLIN INVEST, (1988) 81 (2), 475-479. CODEN: JCINAO. ISSN: 0021-9738.

FS BA; OLD

LA English

AB In autoimmune ***gastritis*** antibodies against a membrane-bound parietal cell antigen of previously unknown function are present in the sera of patients. In this study, a vesicular membrane preparation of porcine ***gastric*** mucosa cells was found to be a potent antigenic source. This preparation blocked > 90% of antibody binding to a lysate of ***gastric*** mucosa cells. The membrane fraction contained H+,K+-ATPase (EC 3.6.1.36) as the major protein, which in sodium dodecyl sulfate-polyacrylamide gel electrophoresis migrated with a weight of 92

kD. After reduction and alkylation, this component was resolved into two bands of similar staining intensity (92 and 88 kD). Immunoblotting analysis showed that sera of patients recognized antigen with pattern identical to the major protein of the vesicular membranes. Protein A-Sepharose beads preincubated with immunoglobulins of five individual patient (but not control) sera were all found to reduce both the H+,K+-ATPase activity and the amount of parietal cell antigen of a preparation of vesicular membranes solubilized in n-octylglucoside. Taken together, the results of this study indicate that the major parietal cell antigen is identical to the acid-producing enzyme, H+K+-ATPase, of the parietal cell.

L2 ANSWER 59 OF 86 MEDLINE

AN 89128963 MEDLINE

DN 89128963

TI An endogenous inhibitor of Na,K-ATPase isolated from human plasma inhibits the acid pump of the stomach.

AU ***Mardh S***

CS Department of Medical and Physiological Chemistry, Uppsala University, Sweden.

SO PROGRESS IN CLINICAL AND BIOLOGICAL RESEARCH, (1988) 268B 417-22. Journal code: PZ5. ISSN: 0361-7742.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 198905

L2 ANSWER 60 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 41

AN 1988:225334 BIOSIS

DN BA85:114569

TI ADRENALINE STIMULATES ACID PRODUCTION IN ISOLATED PIG AND HUMAN PARIETAL CELLS.

AU SONG Y-H; ***MARDH S***; NYREN O; LOOF L

CS DEP. MED. PHYSIOL. CHEM., UPPSALA UNIV. BIOMED. CENT., BOX 575, S-751 23 UPPSALA, SWEDEN.

SO SCAND J GASTROENTEROL, (1988) 23 (1), 35-41. CODEN: SJGRA4. ISSN: 0036-5521.

FS BA; OLD

LA English

AB To investigate the mechanisms of adrenergic stimulation of the parietal cell and to study the possible relationship between the stress hormone adrenaline and duodenal ulcer, the effects of adrenaline and various adrenoceptor agonists and antagonists were investigated in parietal cells isolated from pig stomachs and from endoscopic biopsy specimens taken from the ***gastric*** mucosa of patients. Parietal cell acid production was assayed by the aminopyrine accumulation technique. Adrenaline as the sole drug showed poor or no stimulatory effect but potentiated histamine-stimulated acid production. In the presence of histamine, beta-adrenoceptor agonists caused a stimulation of acid formation with the potency order isoproterenol>adrenaline>noradrenaline. The beta-2-selective antagonist ICI118551 was a more potent inhibitor of acid production than both the beta-1 antagonist practolol and the H2-receptor antagonist cimetidine. Studies of (3H)-dihydroalprenolol (DHA) binding to purified parietal cell membranes showed a protein-concentration-dependent and

specific binding of 2.2 .+- 0.6 pmol DHA/.mu.g. Adrenaline increased
gastric acid production in both pig and human parietal cells, most
likely through a beta-2 receptor on the parietal cell. The adrenaline
stimulatory effect in cells obtained from patients with peptic ulcer was
more pronounced than in cells from non-ulcer patients, which indicates a
possible role of adrenaline in some types of ulcer disease.

L2 ANSWER 61 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 42

AN 1988:133859 BIOSIS

DN BA85:68686

TI ELISA OF PROTON POTASSIUM ATPASE THE PARIETAL CELL ANTIGEN.

AU KARLSSON F A; BURMAN P; LOOF L; OLSSON M; SCHEYNIUS A; ***MARDH S***

CS DEP. INTERN. MED., UNIV. HOSP., S-751 85 UPPSALA, SWED.

SO CLIN EXP IMMUNOL, (1987) 70 (3), 604-610. CODEN: CEXIAL. ISSN: 0009-9104.

FS BA; OLD

LA English

AB Vesicular membranes, purified from porcine ***gastric*** mucosa and rich in H+, K+-ATPase, were used to establish an enzyme-linked immunosorbent assay (ELISA) for determinations of parietal cell autoantibodies. Results obtained with the ELISA correlated well with standard immunofluorescence determinations of parietal cell antibodies based on frozen sections of rat stomach. The ELISA however was about 10-fold more sensitive than the immunofluorescence method and had high specificity. Intra- and interassay coefficients of variation, determined with a patient sera of average positivity, were 5.5% and 18%, respectively. The ELISA detected antibody binding in 23 out of 26 sera from patients with known autoimmune atrophic ***gastritis*** , in five of 25 sera with autoimmune thyroiditis, in five of 20 sera from patients with Graves' disease, in three out of 20 sera from patients with atoxic nodular goitre, in six of 20 sera of patients with primary biliary cirrhosis, in two out of 20 sera of patients with active duodenal ulcer, in two out of 20 sera with detectable antinuclear antibodies, and in one out of 20 sera with detectable rheumatoid factor. Data determined by an ELISA based on a *** gastric *** vesicular membrane preparation of human origin correlated well (r = 0.79, P < 0.001) to those obtained by the standard ELISA based on porcine membrane material. The assay should be well suited for routine determinations of parietal cell antibodies in investigations of autoimmune ***gastritis*** and multiple organ autoimmune endocrinopathies.

L2 ANSWER 62 OF 86 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.DUPLICATE 43

AN 88066330 EMBASE

DN 1988066330

TI Mechanisms of stimulation of acid production in parietal cells isolated from the pig ***gastric*** mucosa.

AU ***Mardh S.***; Song Y.-H.; Carlsson C.; Bjorkman T.

CS Department of Medical and Physiological Chemistry, Biomedical Centre, Uppsala University, S-751 23 Uppsala, Sweden

SO Acta Physiologica Scandinavica, (1987) 131/4 (589-598). ISSN: 0001-6772 CODEN: APSCAX

CY Sweden

DT Journal

FS 002 Physiology

029 Clinical Biochemistry

048 Gastroenterology

037 Drug Literature Index

LA English

SL English

AB Sequential incubations with pronase and collagenase of pig ***gastric*** mucosa resulted in single cell preparations containing 10-20% parietal cells, which could be enriched further to 85-95% purity by density-gradient centrifugation followed by elutriation. Acid production of the isolated cells was measured by means of aminopyrine accumulation in their acid compartments. When small pieces of the mucosa were pretreated for 1 h in the presence of either histamine, pentagastrin or carbachol before preparation of cells, the ability of the subsequently isolated cells to produce acid was increased. In parietal cells isolated from resting (not pretreated) mucosa pentagastrin, carbachol and also adrenaline increased the histamine-stimulated aminopyrine accumulation (50-90% increase). Adrenaline alone had no significant effect on the aminopyrine accumulation. In the presence of 10-4 M histamine the apparent EC50 for adrenaline was 5 x 10-7 M. Adrenaline, histamine, forskolin and isobutylmethylxanthin (IBMX) increased the formation of cAMP in purified parietal cells. The three 'classical' secretagogues histamine, pentagastrin and carbachol, but also IBMX and forskolin, increased the cytosolic free Ca2+ from approximately 1.5 x 10-7 M to 2.2 - 3.5 x 10-7 M but adrenaline and dibutyryl cyclic AMP did not. Thus the present results indicate that there are - in addition to histaminergic H2 receptors specific cholinergic, ***gastrinergic*** and adrenergic receptors on the plasma membrane and that there are separate cAMP and Ca2+-dependent stimulatory pathways in the parietal cell.

L2 ANSWER 63 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1987:435714 BIOSIS

DN BR33:94541

TI ADRENALINE STIMULATES ACID PRODUCTION THROUGH BETA-2 RECEPTORS ON THE PARIETAL CELL.

AU SONG Y; NYREN O; LOOF L; ***MARDH S***

CS DEP. MED., UPPSALA UNIV. BIOMED. CENT., UPPSALA, SWED.

SO TWENTIETH SCANDINAVIAN CONFERENCE ON GASTROENTEROLOGY AND ELEVENTH SCANDINAVIAN MEETING ON GASTROINTESTINAL ENDOSCOPY, TRONDHEIM, NORWAY, JUNE 10-13, 1987. SCAND J GASTROENTEROL SUPPL. (1987) 22 (135), 7. CODEN: SJGSB8. ISSN: 0085-5928.

DT Conference

FS BR; OLD

LA English

L2 ANSWER 64 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 86:652689 SCISEARCH

GA The Genuine Article (R) Number: E8786

TI EFFECTS OF PH ON THE ***GASTRIC*** -ACID PRODUCTION MECHANISM

AU VEGA F V (Reprint); ***MARDH S***

CS UNIV NACL MAR DEL PLATA, DEPT BIOL, MAR DEL PLATA, ARGENTINA

CYA ARGENTINA

SO MEDICINA-BUENOS AIRES, (1986) Vol. 46, No. 5, pp. 494-495.

DT Conference; Journal

FS LIFE; CLIN

LA Spanish

REC No References

L2 ANSWER 65 OF 86 MEDLINE

AN 86182986 MEDLINE

DN 86182986

TI Stimulation of acid formation by histamine, carbachol and pentagastrin in isolated pig parietal cells.

AU Norberg L; Ljungstrom M; Vega F V; ***Mardh S***

SO ACTA PHYSIOLOGICA SCANDINAVICA, (1986 Mar) 126 (3) 385-90. Journal code: 1U4. ISSN: 0001-6772.

CY Sweden

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 198607

AB Free cells were obtained by sequential incubations of pig ***gastric*** mucosa with pronase and collagenase. Approximately 10-15% of the cell population represented parietal cells. Accumulation of aminopyrine (AP) in the acid compartments of parietal cells was used as an index of their acid production. Histamine, carbachol and pentagastrin each independently stimulated aminopyrine accumulation. The initial rate of aminopyrine accumulation, observed after addition of 10(-4) M carbachol or 10(-6) M pentagastrin, were 32% and 10%, respectively, of that observed with 10(-4) M histamine. Steady-state aminopyrine accumulation in the presence of 10(-4) M histamine, 10(-4) M carbachol or 10(-6) M pentagastrin were 6.2 +/- 3.3, 2.6 +/- 0.6 and 3.0 +/- 1.5 pmol AP per 10(4) parietal cells, respectively (mean \pm -SD, n = 5). The EC50 value for histamine was 3.4 +/- 1.4 X 10(-7) M, and for pentagastrin 5.9 +/- 4.2 X 10(-8) M (mean +/-SD, n = 5). The dose-response curve for carbachol was biphasic. A plateau was reached at 10(-5)-10(-4) M carbachol, and for this phase an apparent EC50 of 2.1 +/- 1.4 X 10(-6) M carbachol was calculated (mean +/- SD, n =5). A further increase to 10(-3) M carbachol increased the aminopyrine accumulation. Atropine (10(-6) M) inhibited the response to concentrations up to 10(-4) M carbachol, but was without effect on the histamine- and pentagastrin-stimulation. The H2-receptor antagonist, cimetidine, right-shifted the dose--response curve for histamine. Also, the pentagastrin-stimulated aminopyrine accumulation was inhibited by cimetidine.(ABSTRACT TRUNCATED AT 250 WORDS)

L2 ANSWER 66 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 44

AN 1986:91765 BIOSIS

DN BA81:2181

TI PHOSPHOLIPID ORGANIZATION IN PROTON POTASSIUM ATPASE-CONTAINING MEMBRANES FROM PIG ***GASTRIC*** MUCOSA.

AU OLAISSON H; ***MARDH S***; ARVIDSON G

CS DEP. MED. AND PHYSIOLOGICAL CHEMISTY, BIOMED. CENTER, UNIV. UPPSALA, BOX 575, S-751 23 UPPSALA, SWEDEN.

SO J BIOL CHEM, (1985) 260 (20), 11262-11267.

CODEN: JBCHA3. ISSN: 0021-9258.

FS BA; OLD

LA English

AB The transverse distribution of the phospholipids in vesicular H+-translocating membranes prepared from pig ***gastric*** mucosa was investigated with the aid of phospholipase C, spingomyelinase, and trinitrobenzenesulfonic acid. The major part (80-90%) of the phosphatidylcholine and the phosphatidylethanolamine, 60% of the

phosphatidylserine, and 45% of the sphingomyelin was located on the external, cytoplasmic side of the vesicle membranes. After treatment with phospholipase C the vesicles still behaved as osmometers and appeared as closed vesicles on the electron micrographs. 31P NMR indicated that the phospholipids in untreated vesicles as well as the unhydrolyzed phospholipids in phospholipase C-treated vesicles were arranged in lamellar structures. The 31P NMR spectrum of untreated vesicles to which Pr3+ ions had been added supported the conclusion that the major part of the membrane phospholipids was located on the external surface of the vesicles. A small fraction of the lipids, 3.6 mol %, was found to consist of glycosphingolipids which occurred at a concentration of 52 nmol/mg of protein.

L2 ANSWER 67 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 45

AN 1985:368521 BIOSIS

DN BA80:38513

TI KINETICS OF THE ACID PUMP IN THE STOMACH PROTON TRANSPORT AND HYDROLYSIS OF ATP AND P NITROPHENYLPHOSPHATE BY THE ***GASTRIC*** PROTON POTASSIUM ATPASE.

AU LJUNGSTROM M; ***MARDH S***

CS DEPARTMENT MEDICAL AND PHYSIOLOGICAL CHEMISTRY, BIOMEDICAL CENTER, BOX 575, S-751 23 UPPSALA, SWEDEN.

SO J BIOL CHEM, (1985) 260 (9), 5440-5444. CODEN: JBCHA3. ISSN: 0021-9258.

FS BA; OLD

LA English

AB Hydrolysis of ATP and p-nitrophenyl phosphate by pig H,K-ATPase was investigated. Hydrolysis of ATP was studied at pH 7.4 in vesicles treated with the ionophore nigericin. The kinetic analysis showed negative cooperativity with one high affinity (Km1 = 3 .mu.M) and one low affinity (Km2 = 208 .mu.M) site for ATP. The rate of hydrolysis decreased at 2000 .mu.M ATP indicating a 3rd site for ATP. When the pH was decreased to 6.5 the experimental results followed Michaelis-Menten enzyme kinetics with one low affinity site (Km = 116 .mu.M). Higher concentrations than 750 .mu.M ATP were inhibitory. Proton transport was measured as accumulation of acridine orange in vesicles equilibrated with 150 mM KCl. The transport at various concentrations of ATP in the pH interval from 6.0-8.0 correlated well with the Hill equation with a Hill coefficient between 1.5-1.9. The concentration of ATP resulting in half-maximal transport rate (S0.5) increased from 5 .mu.M at pH 6.0 to 420 .mu.M at pH 8.0. At acidic pH the rate of proton transport decreased at 1000 .mu.M ATP. The K+-stimulated p-nitrophenylphosphatase (pNPPase) activity resulted in a Hill coefficient close to 2 indicating cooperative binding of substrate. The pNPPase was noncompetitively inhibited by ATP and ADP; half-maximal inhibition was obtained at 2 and 100 .mu.M, respectively. Phospholipase C-treated vesicles lost 80% of the pNPPase activity, but the Hill coefficient did not change. These kinetic results are used for a further development of the reaction scheme of the H,K-ATPase.

L2 ANSWER 68 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 85:505937 SCISEARCH

GA The Genuine Article (R) Number: AQQ34

TI PHOSPHOLIPID ORGANIZATION IN H,K-ATPASE-CONTAINING MEMBRANES FROM PIG ***GASTRIC*** -MUCOSA

AU OLAISSON H (Reprint); ***MARDH S***; ARVIDSON G

CS. UNIV UPPSALA, CTR BIOMED, DEPT MED & PHYSIOL CHEM, BOX 575, S-75123 UPPSALA, SWEDEN (Reprint)

CYA SWEDEN

SO JOURNAL OF BIOLOGICAL CHEMISTRY, (1985) Vol. 260, No. 20, pp. 1262-1267.

DT Article; Journal

FS LIFE

LA ENGLISH

REC Reference Count: 24

L2 ANSWER 69 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 46

AN 1986:145323 BIOSIS

DN BA81:55739

TI OMEPRAZOLE CIMETIDINE AND RANITIDINE INHIBITION OF ACID PRODUCTION IN ISOLATED HUMAN PARIETAL CELLS.

AU GUSTAVSSON S; ***MARDH S***; NORBERG L; NYREN O; WOLLERT S

CS DEP. SURG., UNIV. HOSP., S-751 85 UPPSALA, SWED.

SO SCAND J GASTROENTEROL, (1985) 20 (8), 917-921. CODEN: SJGRA4. ISSN: 0036-5521.

FS BA; OLD

LA English

AB The antisecretory properties of omeprazole, cimetidine, and ranitidine were studied in vitro, using human ***gastric*** mucosal cells, which were obtained by sequential pronase and collagenase incubation of small tissue specimens obtained by endoscopic biopsy. Acid production was measured as the accumulation of radioactive aminopyrine in the acid compartments of the parietal cells. Acid production was stimulated via H2-receptors by histamine (10-4 M or 5 .times. 10-6 M) and via intracellular mechanisms by db-cAMP (10-3 M). Omeprazole induced a dose-dependent inhibition of acid production for all stimulators (IC50 = 2 .times. 10-7 M and 3 .times. 10-8 M with high and low concentrations of histamine, respectively, and 5 .times. 10-6 M with db-cAMP). The H2-receptor antagonists dose-dependently inhibited the histamine-stimulated acid production (IC50 for cimetidine = 10-5 M and 10-6 M and for ranitidine = 10-5 M and 2 .times. 10-7 M for high and low concentrations of histamine, respectively). Neither cimetidine nor ranitidine inhibited acid production after intracellular stimulation with db-cAMP. Omeprazole reduced the aminopyrine accumulation stimulated by histamine (10-4 M) already within 5-10 min, whereas cimetidine (10-3 M) and ranitidine (10-4 M) required 20-30 min. The unstimulated level of acid production was also inhibited by omeprazole but not by the H2-receptor antagonists.

L2 ANSWER 70 OF 86 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.DUPLICATE 47

AN 86082312 EMBASE

DN 1986082312

TI Distribution of carbonic anhydrase in cells and membranes isolated from pig ***gastric*** mucosa.

AU Vega F.V.; Olaisson H.; ***Mardh S.***

CS Departmento de Biologia, Facultad de Ciencias Exactas y Naturales, Universidad Nacional de Mar del Plata, Mar del Plata, Argentina

SO Acta Physiologica Scandinavica, (1985) 124/4 (573-579).

CODEN: APSCAX

CY Sweden

DT Journal

FS 002 Physiology

037 Drug Literature Index

048 Gastroenterology

029 Clinical Biochemistry

LA English

AB Mucosal cells were isolated from pig stomach and then fractionated on linear density gradients of Percoll. Different types of cells were identified by their typical staining and morphology. In disrupted cell fractions, hydration of CO2 by carbonic anhydrase was measured by means of pH-state technique. Localization of carbonic anhydrase to certain cell fractions was also studied by histochemical staining. Both parietal cells and carbonic anhydrase were confined to the low and intermediate density fractions of the gradients. Purified membranes from pig ***gastric*** mucosa, which contained the acid pump of the stomach, the H,K-ATPase, also contained a firmly bound carbonic anhydrase of high activity. The enzyme activity in the membranes was inhibited by acetazolamide, furosemide and KSCN. The molecular mass of the carbonic anhydrase was 33 kDA as estimated by its binding of [14C] furosemide followed by polyacrylamide gel electrophoresis. Previous suggestions of a role of carbonic anhydrase as a supplier of H+ in the secretion of acid are supported by its high activity of its localization to the same membrane as the acid pump of the stomach.

L2 ANSWER 71 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 48

AN 1985:332156 BIOSIS

DN BA80:2148

TI A METHOD FOR IN-VITRO STUDIES ON ACID FORMATION IN HUMAN PARIETAL CELLS STIMULATION BY HISTAMINE PENTAGASTRIN AND CARBACHOL.

AU ***MARDH S*** ; NORBERG L; JUNGSTROM M L; WOLLERT S; NYREN O; GUSTAVSSON S

CS DEP. MED. PHYSIOL. CHEM., BIOMED. CENT., UPPSALA UNIV., UPPSALA, SWEDEN.

SO ACTA PHYSIOL SCAND, (1985) 123 (3), 349-354.

CODEN: APSCAX. ISSN: 0001-6772.

FS BA; OLD

LA English

AB Cells were isolated from human ****gastric*** mucosa on a large scale from ****gastric*** resection specimens and on a microscale from endoscopic biopsies by sequential incubations with pronase and collagenase. The accumulation of aminopyrine (AP) was used as an index of acid production in the parietal cells. Basal accumulation was about 0.2 pmol AP/104 parietal cells. Addition of histamine, db [dibutyl]-cAMP, pentagastrin and carbachol increased the aminopyrine accumulation. Maximal accumulation was of the order of 1000-2800% of the control and was obtained after stimulation by 10-4 M histamine and by 10-3 M db-cAMP. Stimulation by pentagastrin and by carbachol reached 200 to 350% of the control. EC50 was 2 x 10-6 M for histamine, 10-8 M for pentagastrin and 4 x 10-6 M for carbachol. Human parietal cells were enriched from a mixture of ****gastric*** mucosal cells by isopycnic centrifugation on density gradients of Percoll. A parietal cell fraction with a purity of 83% was obtained. The density of human parietal cells was estimated to 1.06 g/ml.

L2 ANSWER 72 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 85:504085 SCISEARCH

GA The Genuine Article (R) Number: AQB33

TI PRELIMINARY CHARACTERIZATION OF LOW-DENSITY PLASMA-MEMBRANE FROM BOVINE
GASTRIC -MUCOSA RELATED WITH THE MECHANISM OF ***GASTRIC***
-ACID SECRETION

AU VEGA F V (Reprint); RODRIGUEZ M P; ***MARDH S***

CS UNIV NACL MAR DEL PLATA, FAC CIENCIAS EXACTAS & NAT, DEPT BIOL, RA-7600 MAR DEL PLATA, ARGENTINA, UNIV UPPSALA, CTR BIOMED, DEPT MED & PHYSIOL CHEM, S-75123 UPPSALA, SWEDEN

CYA ARGENTINA; SWEDEN

SO ACTA PHYSIOLOGICA SCANDINAVICA, (1985) Vol. 124, pp. 123.

DT Conference; Journal

FS LIFE

LA ENGLISH

REC Reference Count: 4

L2 ANSWER 73 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 49

AN 1984:133748 BIOSIS

DN BR27:50240

TI PURIFICATION OF PARIETAL CELLS FROM HUMAN ***GASTRIC*** MUCOSA BY CENTRIFUGATION ON A DENSITY GRADIENT OF PERCOLL.

AU ***MARDH S***; NORBERG L; LJUNGSTROM M; WOLLERT S; ADAMI H-O; NYREN O; LOOF L; GUSTAVSSON S

CS DEPARTMENT OF MEDICAL AND PHYSIOL CHEM., UNIVERSITY OF UPPSALA, SWEDEN. SO THE 85TH ANNUAL MEETING OF THE AMERICAN GASTROENTEROLOGICAL ASSOCIATION

HELD IN CONJUNCTION WITH THE AMERICAN ASSOCIATION FOR THE STUDY OF LIVER DISEASE, AND THE GASTROENTEROLOGY STUDY GROUP, NEW ORLEANS, LA., USA, MAY 19-25, 1984. GASTROENTEROLOGY. (1984) 86 (5 PART 2), 1173.

CODEN: GASTAB. ISSN: 0016-5085.

DT Conference

FS BR; OLD

LA English

L2 ANSWER 74 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 50

AN 1985:322910 BIOSIS

DN BA79:102906

TI PREPARATION OF CELLS FROM PIG ***GASTRIC*** MUCOSA ISOLATION OF PARIETAL CELLS BY ISOPYCNIC CENTRIFUGATION ON LINEAR DENSITY GRADIENTS OF PERCOLL.

AU ***MARDH S***; NORBERG L; LJUNGSTROM M; HUMBLE L; BORG T; CARLSSON C

CS DEP. MED. PHYSIOL. CHEM., BIOMED. CENT., UPPSALA UNIV., BOX 575, S-75123 UPPSALA, SWED.

SO ACTA PHYSIOL SCAND, (1984 (RECD 1985)) 122 (4), 607-614. CODEN: APSCAX. ISSN: 0001-6772.

FS BA; OLD

LA English

AB Cells were isolated from pig ***gastric*** mucosa by a combination of mechanical and enzymatic treatment. Isopycnic centrifugation on linear density gradients of Percoll provided a simple and rapid procedure for obtaining highly enriched parietal cells and chief cells. Their densities were 1.06 and 1.10 g/ml, respectively. Isolated mucosal cells attached to Petri dishes coated with fibronectin or collagen. Both parietal cells and chief cells adhered more readily to fibronectin than collagen. Mucosal cells and cells from the Percoll gradients were maintained for up to 1 wk as primary cell cultures. The ability of free parietal cells to produce acid was tested by the 14C-aminopyrine accumulation technique. Maximal accumulation was 2.5 pmol aminopyrine/104 parietal cells after incubation for 45 min at 10-4 M histamine. The EC50 [median effective concentration] for histamine was 5 .times. 10-6 M. The accumulation of aminopyrine at 10-6 M carbachol and 10-7 M pentagastrin were for both secretagogues about

L2 ANSWER 75 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 51

AN 1984:306266 BIOSIS

DN BA78:42746

TI EFFECTS OF PH ON THE INTERACTION OF LIGANDS WITH THE PROTON POTASSIUM ION ATPASE PURIFIED FROM PIG ***GASTRIC*** MUCOSA.

AU LJUNGSTROM M; VEGA F V; ***MARDH S***

CS DEP. MED. PHYSIOL. CHEM., BIOMEDICAL CENT., UPPSAL UNIV., BOX 575, S-751 23 UPPSALA, SWED.

SO BIOCHIM BIOPHYS ACTA, (1984) 769 (1), 220-230. CODEN: BBACAQ. ISSN: 0006-3002.

FS BA; OLD

LA English

AB The effects of K+, Na+ and ATP on the ***gastric*** (H++K+)-ATPase were investigated at various pH. The enzyme was phosphorylated by ATP with pseudo-1st-order rate constant of 3650 min-1 at pH 7.4. This rate constant increased to a maximal value of .apprx. 7900 min-1 when pH was decreased to 6.0. Alkalinization decreased the rate constant. At pH 8.0 it was 1290 min-1. Additions of 5 mM K+ or Na+, did not change the rate constant at acidic pH, while a neutral or alkaline pH a decrease was observed. Dephosphorylation of phosphoenzyme in lyophilized vesicles was dependent on K+, but not on Na+. Alkaline pH increased the rate of dephosphorylation. K+ stimulated the ATPase and p-nitrophenylphosphatase (pNPPase) activities. At high concentrations K+ was inhibitory. Below pH 7.0 Na+ had little or no effect on the ATPase and pnitrophenylphosphatase, while at alkaline pH, Na+ inhibited both activities. The effect of extravesicular pH on transport of H+ was investigated. At pH 6.5 the apparent Km for ATP was 2.7 .mu.M and increased little when K+ was added extravesicularly. At pH 7.5, millimolar concentrations of K+ increased the apparent Km for ATP. Extravesicular K+ and Na+ inhibited the transport of H+. The inhibition was strongest at alkaline pH and only slight at neutral or acidic pH, suggesting a competition between the alkali metal ions and H+ at a common binding site on the cytoplasmic side of the membrane. Two H+-producing reactions as possible candidates as physiological regulators of (H++K+)-ATPase were investigated. Firstly, the hydrolysis of ATP per se, and secondly, the hydration of CO2 and the subsequent formation of H+ and HCO3-. The amount of H+ formed in the ATPase reaction was highest at alkaline pH. The H+/ATP ratio was .apprx. 1 at pH 8.0. When CO2 was added to the reaction medium there was no change in the rate of H+ transport at pH 7.0, but at pH 8.0 the rate increased 4-times upon the addition of 0.4 mM CO2. A possible cooperation is indicated in the production of acid between H+ + K+-ATPase and a carbonic anhydrase associated with the vesicular membrane is indicated.

L2 ANSWER 76 OF 86 LIFESCI COPYRIGHT 2001 CSA

AN 84:46219 LIFESCI

TI Effects of pH on the interaction of ligands with the (H super(+) + K super(+))-ATPase purified from pig ***gastric*** mucosa.

AU Ljungstroem, M.; Vega, F.V.; ***Mardh, S.***

CS Dep. Med. and Physiol. Chem., Biomed. Cent., Uppsala Univ., Box 575, S-751 23 Uppsala, Sweden

SO BIOCHIM. BIOPHYS. ACTA., (1984) vol. 769, no. 1, pp. 220-230.

DT Journal

FS M

LA English

SL English

AB The effects of K super(+), Na super(+) and ATP on the ***gastric*** (H super(+) + K super(+))-ATPase were investigated at various pH. The enzyme was phosphorylated by ATP at pH 7.4. This increased to a maximal when pH was decreased to 6.0. Alkalinization decreased the rate constant. Dephosphorylation of phosphoenzyme in lyophilized vesicles was dependent on K super(+), but not on Na super(+). Alkaline pH increased the rate of dephosphorylation. K super(+) stimulated the ATPase and p-nitrophenylphosphatase activities. The effect of extravesicular pH on transport of H super(+) was investigated. Extravesicular K super(+) and Na super(+) inhibited the transport of H super(+). The inhibition was strongest at alkaline pH and only slight at neutral or acidic pH, suggesting a competition between the alkali metal ions and hydrogen ions at a common binding site on the cytoplasmic side of the membrane. Two H super(+)-producing reactions as possible candidates as physiological regulators of (H super(+) + K super(+))-ATPase were investigated. The results indicate a possible co-operation in the production of acid between the H super(+) + K super(+)-ATPase and a carbonic anhydrase associated with the vesicular membrane.

L2 ANSWER 77 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 52

AN 1984:305455 BIOSIS

DN BA78:41935

TI CHARACTERIZATION OF PROTON TRANSPORTING MEMBRANES FROM RESTING PIG ****GASTRIC*** MUCOSA.

AU LJUNGSTROM M; NORBERG L; OLAISSON H; WERNSTEDT C; VEGA F V; ARVIDSON G; ***MARDH S***

CS DEP. MED. PHYSIOL. CHEM., BIOMED. CENT., UPPSALA UNIV., BOX 575, S-751 23 UPPSALA, SWED.

SO BIOCHIM BIOPHYS ACTA, (1984) 769 (1), 209-219. CODEN: BBACAQ. ISSN: 0006-3002.

FS BA; OLD

LA English

AB Membrane vesicles were purified from resting corpus mucosa of pig stomachs by velocity-sedimentation on a sucrose-Ficoll step gradient. Two vesicular fractions containing the (H++K+)-ATPase were obtained. One fraction was tight towards KCl, the other was leaky. At 21.degree. C maximal (H++ K+)-ATPase activities of 0.8 and 0.4 .mu.mol .cntdot. mg-1 .cntdot. min-1, respectively, were observed in lyophilized vesicles. The vesicles contained a membrane-associated carbonic anhydrase, the activity of which was in 100-fold excess of the maximal ATPase activity. Both vesicular fractions were rich in phosphatidylcholine, phosphatidylethanolamine, sphingomyelin and cholesterol. The characteristics of ion permeability and transport in the tight vesicles were in agreement with corresponding data for vesicles of a tubulovesicular origin in the parietal cell. Measurement of the rate of K+ uptake into the vesicles was based on the ability of K+ to promote H+ transport. The uptake was slow and dependent on the type of anion present. The effectiveness in promoting uptake of K+ by anions was SCN-> NO3-> Cl-.mchgt. HCO3-> SO42-. Uptake of K+ was much more rapid at alkaline pH than at neutral or at acidic pH. Addition of CO2 at alkaline pH strongly stimulated the rate of H+ accumulation in the vesicles. The initial part of this stimulation was sensitive to acetazolamide, an inhibitor of carbonic anhydrase. A model how the (H++

K+)-ATPase and the carbonic anhydrase may cooperate is presented. Membrane vesicles of a tubulovesicular origin can produce acid.

L2 ANSWER 78 OF 86 LIFESCI COPYRIGHT 2001 CSA

AN 84:46215 LIFESCI

TI Characterization of proton-transporting membranes from resting pig ****gastric*** mucosa.

AU Ljungstroem, M.; Norberg, L.; Olaisson, H.; Wernstedt, C.; Vega, F.V.; Arvidson, G.; ***Mardh, S.***

CS Dep. Med. Physiol. Chem., Biomed. Cent., Uppsala University, Box 575, S-751 23 Uppsala, Sweden

SO BIOCHIM. BIOPHYS. ACTA., (1984) vol. 769, no. 1, pp. 209-219.

DT Journal

FS M

LA English

SL English

AB Membrane vesicles were purified from resting corpus mucosa of pig stomachs by velocity-sedimentation on a sucrose-Ficoll step gradient. Two vesicular fractions containing the (H super(+) + K super(+))-ATPase were obtained. One fraction was tight towards KCl, the other was leaky. The vesicles contained a membrane-associated carbonic anhydrase, the activity of which was in 100-fold excess of the maximal ATPase activity. Both vesicular fractions were rich in phosphatidylcholine, phosphatidylethanolamine, sphingomyelin and cholesterol. The characteristics of ion permeability and transport in the tight vesicles were in agreement with corresponding data for vesicles of a tubulovesicular origin in the parietal cell. Measurement of the rate of K super(+) uptake into the vesicles was based on the ability of K super(+) to promote H super(+) transport. The uptake was slow and dependent on the type of anion present. A model how the (H super(+) + K super(+))-ATPase and the carbonic anhydrase may co-operate is presented. It is concluded that membrane vesicles of a tubulovesicular origin can produce acid.

L2 ANSWER 79 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 53

AN 1983:236534 BIOSIS

DN BA75:86534

TI INHIBITION OF ***GASTRIC*** HYDROGEN POTASSIUM ATPASE BY THE SUBSTITUTED BENZIMIDAZOLE PICOPRAZOLE.

AU WALLMARK B; SACHS G; ***MARDH S***; FELLENIUS E

CS AB HASSLE, RES. LAB., S-431 83 MOLNDAL.

SO BIOCHIM BIOPHYS ACTA, (1983) 728 (1), 31-38. CODEN: BBACAQ. ISSN: 0006-3002.

FS BA; OLD

LA English

AB The substituted benzimidazole, picoprazole, inhibited the [hog]

gastric (H++K+)-ATPase in a concentration- and time-dependent manner. Half-maximal inhibition of the (H++K+)-ATPase activity was obtained at .apprx. 2 .cntdot. 10-6 M under standard conditions. In addition to the inhibition of ATPase activity, parallel inhibition of phosphoenzyme formation and the proton transport activity were achieved. Radiolabeled picoprazole was bound to a 100 kDa [kilodalton] peptide; this peptide was shown by phosphorylation experiments to contain the catalytic center of the (H++K+)-ATPase. Studies on the (Na++K+)-ATPase indicated that this enzyme was unaffected by picoprazole. From the data presented and from other pharmacological studies, it is proposed that this compound

inhibits acid secretion at the level of the parietal cell by its ability to inhibit the ***gastric*** proton pump, the (H+ + K+)-ATPase.

L2 ANSWER 80 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 54

AN 1984:16005 BIOSIS

DN BR26:16005

TI EFFECTS OF PHOSPHO LIPASE C ON ***GASTRIC*** VESICLE MEMBRANES CONTAINING HYDROGEN ION POTASSIUM ION ATPASE.

AU OLAISSON H; ***MARDH S***; ARVIDSON G

CS INST. OF MED. AND PHYSIOL. CHEM., BIOMED. CENT., UNIV. OF UPPSALA, BOX 575, S-75123 UPPSALA, SWED.

SO Acta Chem. Scand., Ser. B, (1982 (RECD 1983)) 36 (9), 649-650. CODEN: ACBOCV. ISSN: 0302-4369.

FS BR; OLD

LA English

L2 ANSWER 81 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 55

AN 1983:183275 BIOSIS

DN BA75:33275

TI ATP ADP EXCHANGE ACTIVITY OF ***GASTRIC*** PROTON POTASSIUM ATPASE.

AU RABON E; SACHS G; ***MARDH S***; WALLMARK B

CS LAB. MEMBRANE BIOL., UNIV. ALABAMA BIRMINGHAM, BIRMINGHAM, ALA. 35294.

SO BIOCHIM BIOPHYS ACTA, (1982) 688 (2), 515-524. CODEN: BBACAQ. ISSN: 0006-3002.

FS BA; OLD

LA English

AB The ATP/ADP exchange is shown to be a partial reaction of the [hog] (H++ K+)-ATPase by the absence of measurable nucleoside diphosphokinase activity and the insensitivity of the reaction to P1,P5-di(adenosine-5') pentaphosphate, a myokinase inhibitor. The exchange demonstrates an absolute requirement for Mg2+ and is optimal at an ADP/ATP ratio of 2. The high ATP concentration (K0.5 = 116 .mu.M) required for maximal exchange is interpreted as evidence for the involvement of a low affinity form of nucleotide site. The ATP/ADP exchange is regarded as evidence for an ADP-sensitive form of the phosphoenzyme. In native enzyme, pre-steady state kinetics show that the formation of the phosphoenzyme is partially sensitive to ADP while modification of the enzyme by pretreatment with 5,5'-dithiobis(2-nitrobenzoic acid) (DTNB) in the absence of Mg2+ results in a steady-state phosphoenzyme population, a component of which is ADP sensitive. The ATP/ADP exchange reaction can be either stimulated or inhibited by the presence of K+ as a function of pH and Mg2+.

L2 ANSWER 82 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1982:106645 BIOSIS

DN BR23:36637

TI MONOVALENT CATION DEPENDENT REGULATION OF HYDROGEN ION POTASSIUM ATPASE FROM PIG ***GASTRIC*** MUCOSA.

AU LJUNGSTROM M; ***MARDH S***

CS INST. MED. PHYSIOL. CHEM., UPPSALA, SWED.

SO MEETING OF THE BIOCHEMICAL SOCIETY, MARCH 29-APRIL 3, 1981. BIOCHEM SOC TRANS. (1981) 9 (2), 179P.

CODEN: BCSTB5. ISSN: 0300-5127.

DT Conference

FS BR; OLD

LA English

L2 ANSWER 83 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 82:635282 SCISEARCH

GA The Genuine Article (R) Number: US435

TI MONO-VALENT CATION-DEPENDENT REGULATION OF H&,K&-ATPASE FROM PIG ****GASTRIC*** -MUCOSA

AU LJUNGSTROM M (Reprint); ***MARDH S***

CS UNIV UPPSALA, INST MED & PHYSIOL CHEM, S-75105 UPPSALA, SWEDEN

CYA SWEDEN

SO BIOCHEMICAL SOCIETY TRANSACTIONS, (1981) Vol. 9, No. 2, pp. P179.

DT Conference; Journal

FS LIFE

LA ENGLISH

REC No References

L2 ANSWER 84 OF 86 MEDLINE

AN 80049837 MEDLINE

DN 80049837

TI Phosphorylation and dephosphorylation kinetics of potassium-stimulated ATP phosphohydrolase from hog ***gastric*** mucosa.

AU Wallmark B; ***Mardh S***

SO JOURNAL OF BIOLOGICAL CHEMISTRY, (1979 Dec 10) 254 (23) 11899-902. Journal code: HIV. ISSN: 0021-9258.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 198003

AB Partial reactions of potassium-stimulated ATP phosphohydrolase from hog ***gastric*** mucosa were studied by means of a rapid-mixing apparatus. At 21 degrees C, in the presence of 2 mM MgCl2 and 5 microM [gamma-32P]ATP there was a rapid phosphorylation of the enzyme with a pseudofirst order rate constant of 1400 min-1. Addition of the ATP about 120 ms before the MgCl2 increased this rate constant to 4400 min-1. In the absence of MgCl2 there was no phosphorylation. Addition of 4 or 10 mM KCl to the phosphoenzyme which had been formed in the absence of KCl produced a rapid initial rate of dephosphorylation (k = 2600 and 3200 min-1 respectively). An additional slow component of dephosphorylation was observed when unlabeled ATP was added together with the KCl (k = 700 to 900 min-1). At a 4 mM concentration, KCl stimulated the ATPase activity about 9-fold. At higher concentrations, the activity was reduced in parallel with a reduction of the steady state level of phosphoenzyme. Addition of KCl to the enzyme before the addition of ATP plus MgCl2 resulted in a low rate and extent of phosphorylation. KCl appeared to inhibit the phosphorylation at a level preceeding the E.ATP complex.

L2 ANSWER 85 OF 86 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 79:537048 SCISEARCH

GA The Genuine Article (R) Number: HX412

TI PHOSPHORYLATION AND DEPHOSPHORYLATION KINETICS OF POTASSIUM-STIMULATED ATP PHOSPHOHYDROLASE FROM HOG ***GASTRIC*** -MUCOSA

AU WALLMARK B; ***MARDH S (Reprint)***

CS AB HASSLE, DEPT ANALYT CHEM & BIOCHEM, S-43120 MOLNDAL 1, SWEDEN; UNIV UPPSALA, CTR BIOMED, INST MED & PHYSIOL CHEM, S-75123 UPPSALA, SWEDEN CYA SWEDEN

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SO JOURNAL OF BIOLOGICAL CHEMISTRY, (1979) Vol. 254, No. 23, pp. 1899-1902. DT Article; Journal
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FS LIFE

LA ENGLISH

REC Reference Count: 15

L2 ANSWER 86 OF 86 BIOSIS COPYRIGHT 2001 BIOSIS DUPLICATE 56

AN 1980:54306 BIOSIS

DN BR18:54306

TI REGULATION OF INTERMEDIARY PHOSPHORYLATION OF POTASSIUM ATPASE FROM PIG ***GASTRIC*** MUCOSA BY SODIUM IONS.

AU LJUNGSTROM M; WALLMARK B; ***MARDH S***

CS BIOMED. CENT., INST. MED. PHYSIOL. CHEM., UPPS. UNIV., BOX 575, S-751 23 UPPSALA, SWED.

SO Acta Chem. Scand., Ser. B, (1979 (RECD 1980)) 33 (8), 618-619. CODEN: ACBOCV. ISSN: 0302-4369.

DT Short Communication

FS BR; OLD

LA English

=> e mardh erik/au

- E1 3 MARDH C K/AU
- E2 2 MARDH CARINA KARRMAN/AU
- E3 0 --> MARDH ERIK/AU
- E4 61 MARDH G/AU
- E5 3 MARDH GORAN/AU
- E6 6 MARDH P/AU
- E7 1393 MARDH P A/AU
- E8 1 MARDH P A */AU
- E9 2 MARDH P N/AU
- E10 9 MARDH PER A/AU
- E11 63 MARDH PER ANDERS/AU
- E12 1 MARDH PER ANDERSD/AU
- => s pylori and gastri?
- L3 41957 PYLORI AND GASTRI?
- => s pepsinogen and gastri?
- L4 5756 PEPSINOGEN AND GASTRI?
- => s atpase and gastri?
- L5 8244 ATPASE AND GASTRI?
- => s 13 and 14 and 15
- L6 7 L3 AND L4 AND L5
- => dup rem 16

PROCESSING COMPLETED FOR L6

=> d bib ab 1-

YOU HAVE REQUESTED DATA FROM 5 ANSWERS - CONTINUE? Y/(N):y

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L7 ANSWER 1 OF 5 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.DUPLICATE 1
AN 1998239732 EMBASE
TI Serum antibodies to H+,K+- ***ATPase***, serum ***pepsinogen*** A
   and Helicobacter ***pylori*** in relation to ***gastric*** mucosa
   morphology in patients with low or low-normal concentrations of serum
   cobalamins.
AU Lindgren A.; Burman P.; Kilander A.F.; Nilsson O.; Lindstedt G.
CS Dr. A. Lindgren, Department of Internal Medicine, Boras Central Hospital,
   S-501 82 Boras, Sweden_
SO European Journal of Gastroenterology and Hepatology, (1998) 10/7
   (583-588).
   Refs: 41
   ISSN: 0954-691X CODEN: EJGHES
CY United Kingdom
DT Journal; Article
FS 029 Clinical Biochemistry
  048 Gastroenterology
LA English
SL English
AB Objectives. To compare the diagnostic performance of serum antibodies to
  H+,K+- ***ATPase*** (EC 3.6.1.36), serum ***pepsinogen*** A (EC
  3.4.23.1) and the Schilling test in diagnosing chronic atrophic body
    ***gastritis***; to study the interrelationships between H+,K+-
    ***ATPase*** antibodies, serology for Helicobacter ***pylori***, and
   ***gastric*** morphology. Design. Patients with suspected cobalamin
  deficiency and serum cobalamin < 200 .mu.mol/l were investigated using
  upper gastrointestinal endoscopy, the Schilling test and serum tests for
  H+,K+- ***ATPase*** antibodies, ***pepsinogen*** A, and H.
   ***pylori*** . Setting. The Department of Internal Medicine, Sahlgrenska
  University Hospital, Goteborg, Sweden. Patients. Ninety seven
  consecutively referred patients. Main outcome measures. Sensitivity and
  specificity of assays for serum H+,K+- ***ATPase*** antibodies, serum
   ***pepsinogen*** A, and the Schilling test. Results. Assays of serum
  antibodies to H+,K+- ***ATPase*** and of serum ***pepsinogen*** A
  displayed equal diagnostic sensitivity for atrophic ***gastritis***
  (around 0.90 for the severe forms) and higher than that for the Schilling
  test (0.65). The diagnostic specificity for ***pepsinogen*** A (1.0)
  was higher than for H+,K+- ***ATPase*** antibodies (about 0.80). The
  prevalence of antral ***gastritis*** and positivity for H.
   ***pylori*** antibodies declined with the transition of body
   ***gastritis*** into severe atrophy, while the prevalence of H+,K+-
   ***ATPase*** antibodies increased. Conclusion. ***Pepsinogen*** A is
  preferable to serum H+,K+- ***ATPase*** antibodies in the diagnosis of
   ***gastric*** body mucosal atrophy. The formation of H+,K+-
```

development of ***gastric*** body muscosal atrophy.

ATPase antibodies does not seem to be a primary event in the

AN 97:876684 SCISEARCH

GA The Genuine Article (R) Number: YG753

TI Regulation of ***gastric*** secretion

AU Schubert M L (Reprint)

CS MCGUIRE DEPT VET AFFAIRS MED CTR, DIV GASTROENTEROL, CODE 111N, 1201 BROAD ROCK BLVD, RICHMOND, VA 23249 (Reprint)

CYA USA

SO CURRENT OPINION IN GASTROENTEROLOGY, (NOV 1997) Vol. 13, No. 6, pp. 441-450.

Publisher: RAPID SCIENCE PUBLISHERS, 2-6 BOUNDARY ROW, LONDON, ENGLAND SE1 8NH.

ISSN: 0267-1379.

DT Article; Journal

FS CLIN

LA English

REC Reference Count: 83

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

Gastric secretion is finely regulated by neural, hormonal, and paracrine pathways, During ingestion of a meal, the pathways can be activated by stimuli originating in the brain or stimuli originating in the stomach, such as mechanical stimulation leg, distension) or chemical stimulation leg, protein). The main secretagogues active at the level of the parietal cell are acetylcholine (neurotransmitter), ***gastrin*** (hormone), and histamine (paracrine agent); the main inhibitor is somatostatin (paracrine agent). The release of these four agents by neural, hormonal, and paracrine mechanisms and the interactions among them determine the rate of acid secretion in response to physiological stimuli. The two main intracellular signaling pathways involve cyclic adenosine monophosphate and calcium. A third pathway, involving accumulation of guanosine 3',5'-cyclic monophosphate, is receiving increased attention, Progress has been made in characterizing the intracellular events that occur between generation of second messengers and incorporation of tubulovesicle-rich H+K+- ***ATPase*** into the apical membrane of the parietal cell and exocytosis of ***pepsinogen*** from the chief cell.

L7 ANSWER 3 OF 5 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1998:25516 BIOSIS

DN PREV199800025516

- TI Helicobacter ***pylori*** associated autoantibodies recognize Lewis antigens, and peptide epitopes of ***gastric*** H+, K+- ***ATPase*** and intrinsic factor.
- AU Appelmelk, B. J. (1); Straver, S. (1); Claeys, D.; Faller, G.; Kirchner,
 T.; Negrini, R.; Krakowka, S.; Eaton, K.; Vandenbroucke-Grauls, C. M. J.
 E. (1)
- CS (1) Vrije Univ., Amsterdam Netherlands
- SO Gut, (1997) Vol. 41, No. SUPPL. 1, pp. A17.

Meeting Info.: European Helicobacter Pylori Study Group Xth International Workshop on Gastroduodenal Pathology and Helicobacter Pylori Lisbon, Portugal September 11-14, 1997 European Helicobacter pylori Study Group . ISSN: 0017-5749.

DT Conference

LA English

L7 ANSWER 4 OF 5 SCISEARCH COPYRIGHT 2001 ISI (R) AN 94:715221 SCISEARCH

GA The Genuine Article (R) Number: PQ251

TI POSITIVE CORRELATION BETWEEN H,K-ADENOSINE TRIPHOSPHATASE AUTOANTIBODIES AND HELICOBACTER- ***PYLORI*** ANTIBODIES IN PATIENTS WITH PERNICIOUS-ANEMIA

AU MA J Y; BORCH K; SJOSTRAND S E; JANZON L, MARDH S (Reprint)

CS LINKOPING UNIV, FAC HLTH SCI, DEPT CELL BIOL, S-58185 LINKOPING, SWEDEN (Reprint); LINKOPING UNIV, FAC HLTH SCI, DEPT CELL BIOL, S-58185 LINKOPING, SWEDEN; LINKOPING UNIV HOSP, DEPT MEDICOSURG GASTROENTEROL, S-58185 LINKOPING, SWEDEN; UPPSALA UNIV, CTR BIOMED, DEPT MED & PHYSIOL CHEM, UPPSALA, SWEDEN; AB ASTRA & AB ASTRA ARCUS, SODERTALJE, SWEDEN CYA SWEDEN

SO SCANDINAVIAN JOURNAL OF GASTROENTEROLOGY, (NOV 1994) Vol. 29, No. 11, pp. 961-965.

ISSN: 0036-5521.

DT Article; Journal

FS LIFE; CLIN

LA ENGLISH

REC Reference Count: 30

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB Background: Helicobacter ***pylori*** is a major cause of

gastritis , and the parietal cell H,K-adenosine triphosphatase (

ATPase) is a major autoantigen in autoimmune atrophic corpus

gastritis, which may eventually lead to pernicious anemia and/or neuropathy. Whether the bacterium induces the autoimmune response is unknown. Methods: By means of enzyme-linked immunosorbent assay the occurrence of antibodies against porcine H,K- ***ATPase*** and H.

pylori was determined in sera from 30 patients with pernicious anemia. Results: All sera scored positive against H,K- ***ATPase***, and 25 (83%) scored positive against H. ***pylori***. The titers of antibodies against both antigen preparations inversely correlated with the duration of disease. A possible common epitope in the antigen preparations was tested with a competition assay. There was no indication of a common epitope in either human or porcine H,K- ***ATPase*** and H.

pylori . Conclusions: There was a positive correlation and a high incidence of antibodies against H,K- ***ATPase*** and H. ***pylori*** in sera from patients with pernicious anemia. These antibodies recognized different epitopes.

L7 ANSWER 5 OF 5 MEDLINE

AN 91047803 MEDLINE

DN 91047803

TI Acid and barriers. Current research and future developments for peptic ulcer therapy.

AU Rademaker JW; Hunt RH

CS Division of Gastroenterology, McMaster University Medical Centre, Hamilton, Ontario, Canada...

SO SCANDINAVIAN JOURNAL OF GASTROENTEROLOGY. SUPPLEMENT, (1990) 175 19-26. Ref: 67

Journal code: UCT. ISSN: 0085-5928.

CY Norway

DT Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

LA English

FS Priority Journals

EM 199102

AB Medical therapy for peptic ulcer disease has been targeted at inhibiting acid secretion based on the belief that ulcers occur due to an imbalance between aggressive and protective factors. New antisecretory agents are unlikely to show any dramatic improvement over the success and safety of histamine H2 receptor antagonists or the recently introduced H+K+ ***ATPase*** proton pump antagonist omeprazole. The development of specific muscarinic M3 and ***gastrin*** receptor antagonists will provide useful agents to suppress acid and ***pepsinogen*** secretion by alternative means and may prevent the associated hypergastrinaemia seen with anti-secretory therapy. Enhancement of mucosal defence by site protective agents will be based on a better understanding of the vascular and immune factors involved in maintaining mucosal integrity and the growth factors that regulate wound healing. Molecular techniques are likely to produce the 'model anti-ulcer' agent which will effectively inhibit acid secretion and also enhance wound healing thus providing a cure for this chronic disease.

=> s antibod? and (13 or 14 or 15)

L8 7055 ANTIBOD? AND (L3 OR L4 OR L5)

=> s 18 and (immunoassaY?)

L9 564 L8 AND (IMMUNOASSAY?)

=> dup rem 19

PROCESSING COMPLETED FOR L9 L10 361 DUP REM L9 (203 DUPLICATES REMOVED)

=> s 110 and gastritis

L11 184 L10 AND GASTRITIS

=> d bib ab 1-

YOU HAVE REQUESTED DATA FROM 184 ANSWERS - CONTINUE? Y/(N):y

L11 ANSWER 1 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 2001:104980 BIOSIS

DN PREV200100104980

TI Association of CagA-positive infection with Helicobacter ***pylori***
antibodies of IgA class.

AU Rautelin, Hilpi I. K. (1); Oksanen, Aino M.; Karttunen, Riitta A.; Seppala, Kari M. Y.; Virtamo, Jarmo R. K.; Aromaa, Arpo J.; Kosunen, Timo U.

CS (1) Department of Bacteriology and Immunology, University of Helsinki, 00014, Helsinki: Hilpi Rautelin@Helsinki.fi Finland

SO Annals of Medicine, (December, 2000) Vol. 32, No. 9, pp. 652-656. print. ISSN: 0785-3890.

DT Article

LA English

SL English

AB cagA gene, the best known virulence factor of Helicobacter ***pylori*** , codes for an immunodominant CagA protein. In this study, CagA ***antibodies*** of the IgG class were measured by immunoblot or enzyme ***immunoassay*** in subjects with positive H. ***pylori*** serology, and the presence of CagA ***antibodies*** was compared with that of H. ***pylori*** ***antibodies*** of IgA and IgG classes. Serum samples were available for a total of 1481 subjects, including gastroscopied patients with biopsy-verified H. ***pylori*** infection, smoking men with a normal or low serum ***pepsinogen*** I level indicating atrophic corpus ***gastritis***, and subjects who later developed ***gastric*** cancer and their matched controls. CagA ***antibodies*** were significantly more prevalent among individuals with elevated H. ***pylori*** ***antibody*** titres of the IgA class than in those with IgG ***antibodies*** only, with the exception of a small subgroup of individuals who later developed ***gastric*** cancer. CagA-positive H. ***pylori*** strains seem to induce an immune response with a markedly higher frequency of IgA than what is found in inflammation caused by CagA-negative strains. The presence of serum IgA ***antibodies*** to H. ***pylori*** seems to indicate a higher risk for CagA-positive H. ***pylori*** infection and possibly more severe late sequelae of the disease.

L11 ANSWER 2 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 2000:506124 BIOSIS

DN PREV200000506124

TI Detection of anti-CagA ***antibodies*** in H. ***pylori***
-infected children by an anti-CagA EIA method.

AU Lopez-Brea, M. (1); Martinez, M. J.; Sanz, J. C. (1); Domingo, D. (1); Alacron, T. (1)

CS (1) Hosp. De La Princesa, Madrid Spain

SO Abstracts of the Interscience Conference on Antimicrobial Agents and Chemotherapy, (1999) Vol. 39, pp. 237. cd-rom.

Meeting Info.: 39th Interscience Conference on Antimicrobial Agents and Chemotherapy San Francisco, California, USA September 26-29, 1999 American Society for Microbiology

DT Conference

LA English

SL English

L11 ANSWER 3 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 2000:442192 BIOSIS

DN PREV200000442192

TI Evaluation of blood tests to predict normal ***gastric*** mucosa.

AU Oksanen, A.; Sipponen, P.; Miettinen, A.; Sarna, S.; Rautelin, H. (1)

CS (1) Dept. of Bacteriology and Immunology, University of Helsinki, FIN-00014, Helsinki Finland

SO Scandinavian Journal of Gastroenterology, (August, 2000) Vol. 35, No. 8, pp. 791-795. print. ISSN: 0036-5521.

DT Article

LA English

SL English

AB [Background: To determine the accuracy of blood tests in predicting normal

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***gastric*** mucosa confirmed by histological examination of
    ***gastric*** biopsy specimens. Methods: In total, 207 consecutive
   patients referred for upper endoscopy were included. Two biopsy specimens
   each from the antrum and corpus were assessed histologically for the
   presence of Helicobacter ***pylori***, ***gastritis***, and
   atrophy. Serum samples were studied for H. ***pylori***
    ***antibodies*** by enzyme ***immunoassay*** (Pyloriset EIA-G and
   EIA-A) and by a rapid latex agglutination test (Pyloriset Dry);
    ***pepsinogen*** I was measured by an immunoenzymometric assay
   (Gastroset PGI), ***gastrin*** by radioimmunoassay, and parietal cell
    ***antibodies*** by indirect immunofluorescence. Results: In 101 (49%)
   of 207 patients, the ***gastric*** mucosa on histologic examination
   was normal. In the 63 patients aged 45 years or less, H. ***pylori***
   IgG serology was negative in all 47 patients with normal ***gastric***
   mucosa and none had low serum ***pepsinogen*** I levels. Among 144
   patients over age 45 years, 72 had negative H. ***pylori*** IgG
   serology. Combining the serum ***pepsinogen*** I assay with the
   results of H. ***pylori*** IgG serology, 12 patients with normal
   serology but low serum ***pepsinogen*** I were found. Thus, 60
   patients, 52 of whom showed normal ***gastric*** histology, had normal
   IgG serology and serum ***pepsinogen*** I. In the remaining eight
   patients with normal blood tests, the histologic changes were very mild.
   Conclusions: Although negative H. ***pylori*** IgG serology alone in
   younger patients, and in combination with normal serum ***pepsinogen***
   I levels in older patients, reliably predicted the presence of normal
    ***gastric*** mucosa, gastroscopy is still recommended for patients over
   45 years.
L11 ANSWER 4 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS
AN 2000:182123 BIOSIS
DN PREV200000182123
TI Atrophic ***gastritis*** and Helicobacter -***pylori*** infection
   in outpatients referred for gastroscopy.
AU Oksanen, A.; Sipponen, P.; Karttunen, R.; Miettinen, A.; Veijola, L.;
   Sarna, S.; Rautelin, H. (1)
CS (1) Department of Bacteriology and Immunology, University of Helsinki,
   FIN-00014, Helsinki Finland
SO Gut, (April, 2000) Vol. 46, No. 4, pp. 460-463.
   ISSN: 0017-5749.
DT Article
LA English
SL English
AB Background: Atrophic ***gastritis*** has been shown to be one of the
  long term sequelae of Helicobacter ***pylori*** infection. Aims: To
  determine the prevalence of atrophic ***gastritis*** in outpatients,
  to study the accuracy of serological methods for revealing atrophy, and to
  define the association of H ***pylori*** infection with atrophic
    ***gastritis*** in these patients. Patients/methods: A total of 207
  consecutive outpatients referred for gastroscopy were included. Biopsy
  specimens from the antrum and corpus were assessed histologically
  according to the Sydney system. Serum samples were studied for H
    ***pylori*** IgG and IgA. ***antibodies*** by enzyme
    ***immunoassay*** , CagA ***antibodies*** by immunoblot.
    ***pepsinogen*** I by an immunoenzymometric assay, ***gastrin*** by
  radioimmunoassay, and parietal cell ***antibodies*** by indirect
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immunofluorescence. Results: Histological examination revealed atrophic

***gastritis*** in 52 (25%) of 207 patients. H ***pylori*** and CagA

***antibodies*** were strongly associated with atrophic antral

***gastritis*** but poorly associated with atrophic corpus

***gastritis***. Low serum ***pepsinogen*** I was the most sensitive
and specific indicator of moderate and severe atrophic corpus

***gastritis*** All six patients with moderate atrophic corpus

***gastritis*** had H ***pylori*** infection but eight of 10

patients with severe atrophic corpus had increased parietal cell

***antibodies*** and nine had no signs of H ***pylori*** infection.

Conclusions: Atrophic antral ***gastritis*** was strongly associated

with CagA positive H ***pylori*** infection. Severe atrophic corpus

***gastritis*** was not determined by H ***pylori*** tests but low

serum ***pepsinogen*** I, high ***gastrin***, and parietal cell

***antibodies*** may be valuable in detecting these changes.
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- L11 ANSWER 5 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS
- AN 2000:95018 BIOSIS
- DN PREV200000095018
- TI Evaluation of rapid ***antibody*** tests for the diagnosis of Helicobacter ***pylori*** infection.
- AU Faigel, Douglas O. (1); Magaret, Nathan; Corless, Christopher; Lieberman, David A.; Fennerty, M. Brian
- CS (1) Portland VA Medical Center (P3GI), 3710 US Veterans Hospital Road, Portland, OR, 97201 USA
- SO American Journal of Gastroenterology, (Jan., 2000) Vol. 95, No. 1, pp. 72-77.

ISSN: 0002-9270.

- DT Article
- LA English
- SL English
- AB OBJECTIVE: The aim of this study was to compare the performance characteristics of one serum and four whole blood rapid ***antibody*** tests for Helicobacter ***pylori*** infection. METHODS: A total of 97 outpatients referred for endoscopic evaluation of dyspepsia were included. Antral biopsies were obtained for histology and rapid urease test. Serum was tested with an enzyme-linked ***immunoassay*** (HM-CAP) and a rapid serology test (FlexSure HP). A commercially available 13C-urea breath test was performed. Capillary blood obtained by fingerstick was tested with FlexSure HP, QuickVue, Accustat, and StatSimple ***pylori*** tests. Sensitivity, specificity, and accuracy of each rapid test was calculated relative to a criterion standard of histological
 - ***gastritis*** and at least two of the four following tests positive: identifiable organisms on specially stained slides, rapid urease test, urea breath test, or serum ***immunoassay***. RESULTS: A total of 30 patients (31%) were infected. The FlexSure HP Serum, and FlexSure HP, QuickVue, Accustat, and StatSimple ***pylori*** whole blood tests had sensitivities of 90%, 87%, 83%, 76%, and 90%; specificities of 94%, 90%, 96%, 96%, and 98%, and accuracies of 93%, 88%, 92%, 87%, and 96%, respectively. Sensitivities were not statistically different. StatSimple
 - ***pylori*** was more specific than FlexSure HP whole blood (p < 0.03), and more accurate than FlexSure whole blood (p < 0.024) and Accustat (p < 0.01). Serum ***immunoassay*** was significantly more sensitive (97%) than FlexSure whole blood, QuickVue, and Accustat (p < 0.01), but its specificity (95%) was not statistically different from the rapid tests.

CONCLUSION: Rapid ***antibody*** testing provides an accurate diagnosis of H. ***pylori*** infection. In general, these tests are less sensitive than, but as specific as, standard serology.

- L11 ANSWER 6 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS
- AN 1999:309139 BIOSIS
- DN PREV199900309139
- TI Correlation of serum immunoglobulin G H. ***pylori*** ***antibody*** titers with histologic & endoscopic findings in patients with dyspepsia.
- AU Park, E. M. (1); Shim, S. C. (1); Park, C. Y. (1); Shon, J. I. (1); Jun, W. K. (1); Kim, B. I. (1); Jung, E. S. (1); Lee, S. J. (1); Shin, J. H. (1); Keum, J. S. (1)
- CS (1) Sungkyunkwan Univ Coll of Medicine, Seoul South Korea
- SO Gastroenterology, (April, 1999) Vol. 116, No. 4 PART 2, pp. A277. Meeting Info.: Digestive Disease Week and the 100th Annual Meeting of the American Gastroenterological Association Orlando, Florida, USA May 16-19, 1999 American Gastroenterological Association ISSN: 0016-5085.
- DT Conference
- LA English
- L11 ANSWER 7 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS
- AN 1999:291593 BIOSIS
- DN PREV199900291593
- TI Spontaneous decline of H. ***pylori*** ***antibody*** titres in patients with advanced atrophic ***gastritis***.
- AU Kokkola, Arto (1); Puolakkainen, Pauli (1); Rautelin, Hilpi; Sipponen, Pentti; Haapiainen, Reijo (1); Kivilaakso, Eero (1); Kosunen, Timo
- CS (1) Dept of Surg, Helsinki Univ Ctral Hosp, Helsinki Finland
- SO Gastroenterology, (April, 1999) Vol. 116, No. 4 PART 2, pp. A218. Meeting Info.: Digestive Disease Week and the 100th Annual Meeting of the American Gastroenterological Association Orlando, Florida, USA May 16-19, 1999 American Gastroenterological Association . ISSN: 0016-5085.
- DT Conference
- LA English
- L11 ANSWER 8 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS
- AN 1999:205896 BIOSIS
- DN PREV199900205896
- TI ***Antibody*** against Helicobacter ***pylori*** CagA and VacA and the risk for ***gastric*** cancer.
- AU Yamaoka, Y. (1); Kodama, T.; Kashima, K.; Graham, D. Y.
- CS (1) Department of Medicine, Veterans Affairs Medical Center (111D), 2002 Holcombe Blvd, Houston, TX, 77030 USA
- SO Journal of Clinical Pathology (London), (March, 1999) Vol. 52, No. 3, pp. 215-218.
 - ISSN: 0021-9746.
- DT Article
- LA English
- SL English
- AB Aim-Helicobacter ***pylori*** is associated with ***gastric*** cancer. Our aim was to investigate whether CagA or VacA seropositivity provides additional risk for ***gastric*** cancer. Methods-Sera from 110 ***gastric*** cancer patients were sex and aged matched with

asymptomatic controls. H ***pylori*** status was determined by IgG enzyme ***immunoassay*** (HM-CAP EIA); CagA status was assessed by enzyme linked immunosorbent assay (ELISA) (OraVax) and immunoblotting (Chiron), and VacA status by immunoblotting using recombinant proteins as antigens. Results-H ***pylori*** infection was associated with an increased risk of ***gastric*** cancer (odds ratio (OR) = 2.19, 95% confidence interval 1.17 to 4.1). Subgroup analysis showed a significant association with intestinal type (OR = 2.94, 1.35 to 6.41), distal type (OR = 2.97, 1.39 to 6.33), early ***gastric*** cancer (OR = 3.74, 1.54 to 9.06), and age ltoreq 55 years (OR = 8.33, 2.04 to 34.08), but not with diffuse type (OR = 0.83), proximal type (OR = 1.0), advanced ***gastric*** cancer (OR = 1.13), or age > 55 years (OR = 1.40). Serum CagA IgG and VacA ***antibody*** positivity was present in similar proportions in patients with and without cancer, with no significant differences in histological classification, clinical stage, or location (p > 0.3). Conclusions-H ***pylori*** infection causes chronic ***gastritis*** and is associated with the development of ***gastric*** cancer. Neither CagA nor VacA seropositivity added additional information or stratification.

L11 ANSWER 9 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1999:156989 BIOSIS

DN PREV199900156989

TI Detection of Helicobacter ***pylori*** in ***gastric*** biopsies by PCR: Correlation with conventional methods.

AU Polidorou, F. (1); Vavatsi, N.; Pavlitou, K. (1); Malaka, E. (1)

CS (1) Dep. Microbiol., Gen. Hosp. "Agios Dimitrios", Thessaloniki Greece

SO Deltion Ellenikes Mikrobiologikes Etaireias, (May-June, 1998) Vol. 43, No. 3, pp. 276-279.

ISSN: 0438-9573.

DT Article

LA Greek

SL Greek; English

AB The aim of the study was to evaluate the role of PCR technique in the detection of Helicobacter ***pylori*** (HP) in ***gastric*** biopsies. Antral biopsies from 27 patients with chronic ***gastritis*** and peptic ulcer were examined by means of a rapid urease test, Giernsa and haematoxylin-eosin stains and PCR. Blood samples were tested for IgG ***antibodies*** against HP using an enzyme ***immunoassay***. After the extraction of DNA, nested PCR was performed by using 2 pairs of primers for the amplification of the urease A gene of HP. The final PCR product was 361bp and it was recognised by electrophoresis on a 2% agarose gel. Nineteen out of 27 patients were positive for HP using PCR. The staining method and the ***immunoassay*** detected HP in 18 patients while CLOtest was positive in 17 patients. In one patient HP was detected only by PCR.

L11 ANSWER 10 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1999:60264 BIOSIS

DN PREV199900060264

TI Severe ***gastritis*** in guinea-pigs infected with Helicobacter ***pylori***.

AU Sturegard, E.; Sjunnesson, H.; Ho, B.; Willen, R.; Aleljung, P.; Ng, H. C.; Wadstrom, T. (1)

CS (1) Dep. Med. Microbiol., Lund Univ., Lund Sweden

SO Journal of Medical Microbiology, (Dec., 1998) Vol. 47, No. 12, pp. 1123-1129.

ISSN: 0022-2615.

DT Article

LA English

AB An appropriate animal model is essential to study Helicobacter

pylori infection. The aim of this study was to investigate if H.

pylori can colonise the guinea-pig stomach and whether the infection causes

gastritis and a serological response similar to that observed in man. Guinea-pigs were infected either with fresh H.

pylori isolates from human

gastric biopsies or with a guinea-pig passaged strain. When the animals were killed, 3 and 7 weeks after inoculation, samples were taken for culture, histopathology and serology. H.

pylori was cultured from 22 of 29 challenged animals. All culture-positive animals exhibited a specific immune response

against H. ***pylori*** antigens in Western blotting and ***gastritis*** in histopathological examination. ***Antibody*** titres in enzyme ***immunoassay*** were elevated among animals challenged with H. ***pylori***. The inflammatory response was graded as severe in most animals and consisted of both polymorphonuclear leucocytes and lymphocytes. Erosion of the ***gastric*** epithelium was found in infected animals. These results suggest that the guinea-pig is suitable for studying H. ***pylori*** -associated diseases. Moreover, guinea-pigs are probably more similar to man than any other small laboratory animal as regards ***gastric*** anatomy and physiology.

L11 ANSWER 11 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1999:39237 BIOSIS

DN PREV199900039237

TI Serodiagnosis of Helicobacter ***pylori*** infection in Korean patients using enzyme-linked immunosorbent assay.

AU Kim, S. Y.; Ahn, J. S.; Ha, Y. J.; Doh, H. J.; Jang, M. H.; Chung, S. L.; Park, H. J. (1)

CS (1) MOGAM Biotechnology Res. Inst., 341 Pojung-Ri, Koosung-Myon, Yongin-City, Kyonggi-Do 449-910 South Korea

SO Journal of Immunoassay, (Nov., 1998) Vol. 19, No. 4, pp. 251-270. ISSN: 0197-1522.

DT Article

LA English

AB Helicobacter ***pylori*** (H. ***pylori***) is a gram-negative spiral bacteria that are associated with ***gastritis***, peptic ulcer and ***gastric*** cancer. We have developed enzyme-linked immunosorbent assay (ELISA) that detects serum anti-H. ***pylori*** immunoglobulin G ***antibodies*** using H. ***pylori*** strains isolated from Korean patients. To assess the sensitivity and specificity of our assay system with different commercial kits, serum samples from 249 Korean patients with a variety of gastrointestinal diseases were tested. Among 249 Korean patients, 178 (71.5%) were positive in culture and/or urease test. The sensitivity and specificity between our assay system and four other commercial kits (Bio-Rad, DAKO, ROCHE, and IPR) were as follows: 97.8% and 92%, 94.3% and 53%, 56.5% and 92%, 83.3% and 96%, 58.2% and 92%, respectively. All sera showing discordant ***immunoassay*** results between different ELISA kits were confirmed by immunoblot analysis. These results indicate that our assay system showed a highly

accurate and reliable results in diagnosis of H. ***pylori*** infection in Korean patients.

L11 ANSWER 12 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1998:445623 BIOSIS

DN PREV199800445623

TI Helicobacter ***pylori*** -positive ***gastritis*** in pediatric patients with chronic inflammatory bowel disease.

AU Kolho, Kaija-Leera; Rautelin, Hilpi; Lindahl, Harry; Savilahti, Erkki (1)
 CS (1) Hosp. Children Adolescents, Stenbackinkatu 11, FIN-00290 Helsinki Finland

SO JPGN, (Sept., 1998) Vol. 27, No. 3, pp. 292-295.

DT Article

LA English

AB Background: ***Gastritis*** is a common finding in patients with inflammatory bowel disease. However, the association of ***gastritis*** with Helicobacter ***pylori*** is unclear in these patients. Methods: The prevalence of ***antibodies*** for H. ***pylori*** in serum was determined in 47 pediatric patients with inflammatory bowel disease (19 with Crohn's disease, 21 with ulcerative colitis, and 7 with unclassified disease). H. ***pylori*** ***antibodies*** of the IgG and IgA classes were measured by enzyme ***immunoassay*** in 24 patients at the time of diagnosis of inflammatory bowel disease and in 23 more patients during the follow-up of inflammatory bowel disease (mean follow-up, 3.5 years; range 1-10 years). Esophagogastroduodenoscopy was performed on 40 patients during the examination for inflammatory bowel disease. Results: In contrast to earlier findings, no patient was determined to be positive for H. ***pylori***, either in serologic or histologic examination. This negative finding was unexpected, because it is known that approximately 10% of asymptomatic Finnish children have ***antibodies*** for H. ***pylori*** in serum and approximately 10% of analyses of specimens obtained in ***gastric*** antral biopsies obtained at the Hospital for Children and Adolescents, Helsinki, Finland, are positive for H. ***pylori*** . Conclusions: Permanent colonization of the stomach with H. ***pylori*** is unusual in children with inflammatory bowel disease.

L11 ANSWER 13 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1998:304689 BIOSIS

DN PREV199800304689

TI Positive result by serology indicates active Helicobacter ***pylori*** infection in patients with atrophic ***gastritis***.

AU Kokkola, Arto (1); Rautelin, Hilpi; Puolakkainen, Pauli; Sipponen, Pentti; Farkkila, Martti; Haapiainen, Reijo; Kosunen, Timo U.

CS (1) Second Dep. Surgery, Helsinki Univ. Central Hosp., Haartmaninkatu 4, FI-00290 Helsinki Finland

SO Journal of Clinical Microbiology, (June, 1998) Vol. 36, No. 6, pp. 1808-1810.

ISSN: 0095-1137.

DT Article

LA English

AB Patients with atrophic corpus ***gastritis*** and elevated
Helicobacter ***pylori*** ***antibody*** titers but 130 -urea
breath test (13C-UBT) and histology results negative for H. ***pylori***
were randomized into eradication therapy or follow-up only.

Antibody levels decreased significantly in six out of seven patients in the eradication group, while in the follow-up group, the titers declined in only one out of eight patients. In patients with atrophic corpus ****gastritis***, positive serology results may indicate an ongoing infection in spite of negative 13C-UBT and histology results.

L11 ANSWER 14 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1998:229535 BIOSIS

DN PREV199800229535

TI Evaluation of pyloriset screen, a rapid whole-blood diagnostic test for Helicobacter ***pylori*** infection.

AU Oksanen, Aino; Veijola, Lea; Sipponen, Pentti; Schauman, Knut-Olof; Rautelin, Hilpi (1)

CS (1) Dep. Bacteriol. Immunol., P.O. Box 21, Univ. Helsinki, Helsinki FIN-00014 Finland

SO Journal of Clinical Microbiology, (April, 1998) Vol. 36, No. 4, pp. 955-957.

ISSN: 0095-1137.

DT Article

LA English

AB Helicobacter ***pylori*** infection can be detected by several invasive tests based on gastroscopy and by noninvasive methods such as serologic assays. Noninvasive tests can be used not only in addition to invasive tests but also by themselves to screen for H. ***pylori*** infection in patients who are not in urgent need of endoscopy. Lately, rapid qualitative serologic tests have been developed. In the present study, the accuracy of a novel rapid whole-blood test, Pyloriset Screen, detecting immunoglobulin G (IgG) and IgA ***antibodies*** against H. ***pylori*** was evaluated. A total of 207 consecutive adult outpatients referred for upper endoscopy were enrolled. ***Gastric*** biopsy specimens were taken from the antrum and corpus for histologic examination and rapid urease testing. Cultures were available for 113 patients. Serum samples collected from all patients were tested for H. ***pylori*** ***antibodies*** by two enzyme ***immunoassays*** (EIAs) (Pyloriset EIA and an in-house EIA), a rapid latex agglutination test (Pyloriset Dry), and Pyloriset Screen. Patients were considered H. ***pylori*** positive if helicobacters were seen on histologic examination (77 patients) or, if in combination with histologically verified (although helicobacter-negative) ***gastritis***, their IgG ***antibody*** titers were elevated in the two EIAs (five patients). The Pyloriset Screen test had a sensitivity of 95%, a specificity of 94%, a positive predictive value of 91%, and a negative predictive value of 97%. Among 63 patients under the age of 45 years, the Pyloriset Screen test did not miss a single H. ***pylori*** diagnosis, and only 1 patient had a false-positive result. Pyloriset Screen could be used reliably to screen for H. ***pylori*** infection.

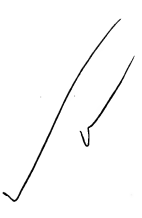
L11 ANSWER 15 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1998:167615 BIOSIS

DN PREV199800167615

TI A new histological procedure for re-evaluation of the serologial test for Helicobacter ***pylori***.

AU Misawa, K. (1); Kumagai, T.; Shimizu, T.; Furihata, K.; Ota, H.; Akamatsu, T.; Katsuyama, T.



CS (1) Dep. Lab. Med., Shinshu Univ. Med., Asahi 3-1-1 Matsumoto 390 Japan
 SO European Journal of Clinical Microbiology & Infectious Diseases, (Jan., 1998) Vol. 17, No. 1, pp. 14-19.
 ISSN: 0934-9723.

DT Article

LA English

AB To re-evaluate the accuracy of the serological test for Helicobacter ***pylori***, fixation of biopsy specimens with Carnoy's solution (preserving the mucous layer in tissue preparations) followed by immunohistochemical staining (a new histological procedure) was used as the reference histological method instead of 10% formalin fixation followed by hematoxylin-eosin staining (the conventional histological procedure). Biopsy specimens (antrum and body) from 114 patients with ***gastritis*** (including non-ulcer dyspepsia) or ***gastric*** and/or duodenal ulcers were obtained by endoscopy and used for both bacteriological culture and histological examination. Serum samples were taken from all patients at the time of endoscopy. The serum levels of specific IgG and IgA ***antibodies*** for Helicobacter ***pylori*** were measured by commercial enzyme ***immunoassay*** kits. The reliability of the IgG and IgA measurements was evaluated by analyzing receiver operating characteristic curves obtained using the two histological procedures. With the conventional histological procedure as the reference, the sensitivity and specificity levels of the serological test were 87.2% and 82.1%, respectively. With the new histological procedure as reference, sensitivity and specificity were 94% and 96.7%, respectively. The insufficient accuracy reported for the serological test could be due to false-positive or false-negative results obtained when the conventional histological procedure is used as the reference. The new histological procedure used here revealed that the serological test for Helicobacter ***pylori*** is more reliable than previously thought.

L11 ANSWER 16 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1997:501874 BIOSIS

DN PREV199799801077

TI Reactivities of Lewis antigen monoclonal ***antibodies*** with the lipopolysaccharides of Helicobacter ***pylori*** strains isolated from patients with gastroduodenal diseases in Japan.

AU Amano, Ken-Ichi (1), Hayashi, Shyunji; Kubota, Toru; Fujii, Nobuhiro; Yokota, Shin-Ichi

CS (1) Central Res. Lab., Akita Univ. Sch. Med., 1-1-1 Hondo, Akita, Akita 010 Japan

SO Clinical and Diagnostic Laboratory Immunology, (1997) Vol. 4, No. 5, pp. 540-544.

ISSN: 1071-412X.

DT Article

LA English

AB We have examined the reactivity of monoclonal ***antibodies*** (Mabs) specific for Lewis antigens (Le-x, Le-y, Le-a, and Le-b) with Helicobacter ***pylori*** lipopolysaccharides (LPS) by immunoblot analysis and enzyme-linked immunosorbent assay (ELISA). Sixty-eight strains of H. ***pylori*** were isolated from patients with chronic ***gastritis***, ***gastric*** and duodenal ulcers, and ***gastric*** cancer in Japan. The cells were treated with proteinase K, and the resulting fractions were used as a source of LPS for the ***immunoassays*** In the immunoblot analysis, 28 isolates (41%) and 29 isolates (42%) reacted

with anti-Le-x and anti-Le-y MAbs, respectively, while 4 isolates (6%) and 7 isolates (10%) reacted with anti-Le-b and anti-Le-b MAbs. On the other hand, in ELISA, the number of isolates that reacted with anti-Le-x MAbs fell significantly to 21 isolates (30%) but the number of isolates that reacted with the other anti-Lewis antigen MAbs remained relatively unchanged. These data show that the immunoblotting technique is more sensitive than the ELISA technique for the detection of immunocomplexes of anti-Le-x MAbs and components of H. ***pylori*** LPS. Furthermore, human serum was found to react with the synthetic Lewis antigens regardless of the status of the individual's H. ***pylori*** infection. This means that humans may naturally possess ***antibodies*** against Lewis antigens in the absence of H. ***pylori*** infection.

L11 ANSWER 17 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1997:35048 BIOSIS

DN PREV199799341451

- TI Seroprevalence of Helicobacter ***pylori*** in south Sweden and Iceland
- AU Bergenzaun, P.; Kristinsson, K. G.; Thjodleifsson, B.; Sigvaldadottir, E.; Molstad, S.; Held, M.; Wadstrom, T. (1)
- CS (1) Dep. Medical Microbiol., Lund University, Solvegatan 23, S-22362 Lund Sweden
- SO Scandinavian Journal of Gastroenterology, (1996) Vol. 31, No. 12, pp. 1157-1161.

ISSN: 0036-5521.

DT Article

LA English

AB Background: Seroepidemiologic studies on the prevalence of Helicobacter

pylori infection have been reported from several European
countries but not from Sweden or Iceland. Methods: Serum samples were
collected from 443 persons in Sweden and 387 persons in Iceland. All the
830 sera were tested with the same enzyme ***immunoassay*** test with
an acid glycine extract of H. ***pylori*** surface proteins as
antigen. Results: The ***antibody*** levels were low in the young age
groups in both Sweden and Iceland, with increasing levels with age.
Conclusions: The results are consistent with previous studies from other
comparable countries, but with important differences. The prevalence was
lower in Sweden than in other, previously studied, Western European
countries, but, on the other hand, the prevalence was slightly higher in
Iceland.

L11 ANSWER 18 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1996:269657 BIOSIS

DN PREV199698825786

- TI Evaluation of three commercial enzyme ***immunoassays*** compared with the 13C urea breath test for detection of Helicobacter ***pylori*** infection.
- AU Marchildon, P. A. (1); Ciota, L. M.; Zamaniyan, F. Z.; Peacock, J. S.; Graham, D. Y.
- CS (1) Enteric Products Inc., 25 E. Loop Rd., Stony Brook, NY 11790 USA
- SO Journal of Clinical Microbiology, (1996) Vol. 34, No. 5, pp. 1147-1152. ISSN: 0095-1137.

DT Article

LA English

AB The diagnostic significance of the serological detection of

antibodies to Helicobacter ***pylori*** has been established by numerous investigators. Reports of the clinical reliabilities of commercial enzyme ***immunoassay*** (EIA) kits available for this purpose vary as a result of the different H. ***pylori*** antigen sources and reference methods used. The 13C urea breath test (UBT) has been shown to be an extremely accurate and reliable method of detecting H. ***pylori*** infection. We used the 13C urea breath test as the confirmatory method for H. ***pylori*** status to evaluate three commercially available EIA kits designed to detect immunoglobulin G ***antibodies*** to H. ***pylori*** . These kits were the HM-CAP EIA kit (Enteric Products, Inc.), the ***PYLORI*** STAT EIA kit (BioWhittaker, Inc.), and the G.A.P. kit (Bio-Rad Laboratories/Biomerica, Inc.). The evaluations were performed in a double-blind manner with samples from 473 clinically characterized patients. This group included patients with symptomatic gastrointestinal disorders as well as nonsymptomatic volunteers. The sensitivities of the kits were as follows: HM-CAP, 98.4%; ***PYLORI*** STAT, 99.2%; and GA.P., 100%. The specificities were as follows: HM-CAP, 96.4%; ***PYLORI*** STAT, 90.1%; and G.A.P., 26.0%. Although the HM-CAP and ***PYLORI*** STAT kits performed comparably, the G.A.P. test yielded significantly more false-positive results and an unacceptably high number of indeterminate results.

AN 1996:221204 BIOSIS DN PREV199698777333 TI High frequency of Helicobacter negative ***gastritis*** in patients with Crohn's disease. AU Halme, L. (1); Karkkainen, P.; Rautelin, H.; Kosunen, T. U.; Sipponen, P. CS (1) Dep. Surg., Helsinki Univ. Hosp., Kasarmikatu 11-13, FIN-00130 Helsinki Finland SO Gut, (1996) Vol. 38, No. 3, pp. 379-383. ISSN: 0017-5749. DT Article LA English AB The frequency of ***gastric*** Crohn's disease has been considered low. This study was undertaken to determine the prevalence of chronic ***gastritis*** and Helicobacter ***pylori*** infection in patients with Crohn's disease. Oesophagogastroduodenoscopy was performed on 62 consecutive patients suffering from ileocolonic Crohn's disease. Biopsy specimens from the antrum and corpus were processed for both histological and bacteriological examinations. H. ***pylori*** ***antibodies*** of IgG and IgA classes were measured in serum samples by enzyme ***immunoassay*** . Six patients (9.7%) were infected with H. ***pylori***, as shown by histology, and in five of them the infection was also verified by serology. Twenty one patients (32%) had chronic H. ***pylori*** negative ***gastritis*** (negative by both histology and serology) and one of them also had atrophy in the antrum and corpus. Granulomas were found in four patients. The characteristic appearance of H. ***pylori*** negative ***gastritis*** was focal and mostly mild inflammation resembling the inflammatory changes seen in the gut in Crohn's disease. Patients with H. ***pylori*** negative chronic ***gastritis*** had a significantly more active disease in their gut than those with normal ***gastric*** mucosa (p lt 0.01). It is concluded that H. ***pylori*** positive ***gastritis*** is rare,

L11 ANSWER 19 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

while H. ***pylori*** negative ***gastritis*** is relatively common in patients with Crohn's disease. H. ***pylori*** negative 'Crohn's ***gastritis*** ' seems to be associated with active Crohn's disease.

L11 ANSWER 20 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1995:550325 BIOSIS

DN PREV199698564625

TI ***Antibody*** titres in Helicobacter ***pylori*** infection: Implications in the follow-up of antimicrobial therapy.

AU Kosunen, Timo U.

CS Dep. Bacteriology Immunology, P.O. Box 21, FIN-00014 Univ. Helsinki, Helsinki Finland

SO Annals of Medicine, (1995) Vol. 27, No. 5, pp. 605-607. ISSN: 0785-3890.

DT Article

LA English

AB Regular presence and persistence of specific serum ***antibodies*** in Helicobacter ***pylori*** infection gives an excellent tool for diagnostic work. Eradication of the infection leads to gradual disappearance of the ***gastritis*** and decrease of specific serum ***antibodies***. The fall of IgG and IgA ***antibody*** titres can be followed with quantititative enzyme ***immunoassays***. Success in eradication is reflected in 40-50% decrease of ***antibody*** titres within 5-6 months. The decrease continues and most patients have normal titres within 2 years. Serology offers a cheap and convenient way to follow the treated patients and makes most follow-up endoscopies unnecessary.

L11 ANSWER 21 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1995:439431 BIOSIS

DN PREV199598453731

TI Helicobacter ***pylori*** infection in recurrent abdominal pain in childhood: Comparison of diagnostic tests and therapy.

AU Chong, Sonny K. F. (1); Lou, Qinyuan; Asnicar, Mark A.; Zimmerman, Sarah E.; Croffie, Joseph M.; Lee, Chao-Hung; Fitzgerald, Joseph F.

CS (1) Dep. of Pediatrics, James Whitcomb Riley Hospital for Children, Indiana Univ. School of Med., 702 Barnhill Dr., Room 2728, Indianapolis, IN 46202-5225 USA

SO Pediatrics, (1995) Vol. 96, No. 2 PART 1, pp. 211-215. ISSN: 0031-4005.

DT Article

LA English

AB Objective: To determine the role of Helicobacter ***pylori*** infection in children with recurrent abdominal pain and the usefulness of serologic tests in screening H. ***pylori*** infection and monitoring treatment of H. ***pylori*** -associated ***gastritis***. Methods: During a 3 year period, we investigated the presence of serum immunoglobulin G (IgG) ***antibody*** to H. ***pylori*** in 456 children using the high-molecular-weight cell-associated protein H. ***pylori*** enzyme ***immunoassay*** kit. Among the 456 children studied, 218 (age range, 3 to 18 years; mean age, 9.5 years) had symptoms of recurrent abdominal pain (RAP syndrome) with or without vomiting, and the remaining 238 (age range, 3 to 18 years; mean age, 9.8 years) had no RAP (non-RAP syndrome). We performed upper gastrointestinal endoscopy on

111 consecutive children of the 218 with RAP syndrome and obtained mucosal biopsies for culture, histologic analysis, CLO test (Delta West, Perth, Australia), and H. ***pylori*** detection by polymerase chain reaction. Results: Thirty-eight (17.4%) of 218 children in the RAP group and 25 (10.5%) of 238 children in the non-RAP group were seropositive for H. ***pylori***. Of the 111 children endoscoped, 95 were found to be negative, and 12 were positive by all five assays. Specimens from 2 children were negative by culture and the CLO test but positive by the other three assays. Specimens from 1 child were negative by histologic analysis but positive by all other tests. The remaining child was positive for anti-H. ***pylori*** IgG but negative by all of the other four assays. Upper gastrointestinal endoscopy detected 14 children with peptic ulcer disease (9 duodenal ulcer and 5 ***gastric*** ulcer) and 12 with antral nodular ***gastritis*** . Only 4 of the 14 diagnosed with peptic ulcer were H. ***pylori*** positive by all five assays, whereas all 12 children with antral nodular ***gastritis*** were H. ***pylori*** positive. Nine of the 12 H. ***pylori*** -positive children were treated with a combination of bismuth subsalicylate, amoxicillin, and metronidazole for 2 weeks. Sera obtained at 2, 4, and 6 months after treatment from all 9 children showed a decrease in anti-H. ***pylori*** IgG titer. Three H. ***pylori*** -infected children who did not receive any treatment served as control children, and their IgG levels remained elevated or increased over time. Conclusion: The results from our study indicate that screening for the serum IgG ***antibody*** to H. ***pylori*** is a practical method for diagnosing H. ***pylori*** infection in children, and that serial measurements of the H. ***pylori*** IgG ***antibody*** are useful for monitoring treatment of H. ***pylori*** because of its high sensitivity and ease of performance. Only 4 of the 14 children diagnosed with peptic ulcer disease were confirmed to be infected with H. ***pylori***, whereas all 12 children with antral nodular ***gastritis*** were found to be infected by H. ***pylori*** . These observations suggest that H. ***pylori*** infection is more frequently associated with ***gastritis*** than with peptic ulcer disease in children, and that H. ***pylori*** ***gastritis*** is a cause of RAP syndrome in children.

L11 ANSWER 22 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1995:245551 BIOSIS

DN PREV199598259851

TI Evaluation of an Enzyme ***Immunoassay*** (Helisal) for Detection of Salivary ***Antibody*** to Helicobacter ***pylori***.

AU Lin, E. (1); Simor, A. E.; Pearen, S.; Saibil, F.; Cohen, L.; Hung, S.; Donhoffer, H. A.; Louie, M.

CS (1) Sunnybrook Health Sci. Cent., Univ. Toronto, Toronto, ON Canada

SO Gastroenterology, (1995) Vol. 108, No. 4 SUPPL., pp. A150. Meeting Info.: 95th Annual Meeting of the American Gastroenterological Association and Digestive Disease Week San Diego, California, USA May 14-17, 1995 ISSN: 0016-5085.

DT Conference

LA English

L11 ANSWER 23 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1995:228024 BIOSIS

DN PREV199598242324

TI Helicobacter ***pylori*** infection in rural settlements (Kibbutzim) in Israel.

AU Gilboa, S. (1); Gabay, G.; Zamir, D.; Zeev, A.; Novis, B.

CS (1) Epidemiol. Unit, Meir Hosp., Kfar-Sava 44281 Israel

SO International Journal of Epidemiology, (1995) Vol. 24, No. 1, pp. 232-237. ISSN: 0300-5771.

DT Article

LA English

AB Background: Helicobacter ***pylori*** (HP) is accepted as a major cause of type B ***gastritis***, which is strongly associated with peptic ulcer disease. Epidemiological studies have indicated a correlation of HP infection and socioeconomic class. Methods: To determine the prevalence of HP infection and to evaluate symptoms and risk factors associated with HP infection in a rural population, 377 asymptomatic individuals were studied out of a random sample of 453 people. Subjects were randomly chosen in a ratio of 1:4 of all the adults over 30 years, living in eight communal settlements in Israel. Blood samples were taken and subjects answered a questionnaire in which sociodemographic information clinical gastrointestinal background and the use of medication were included. A sensitive enzyme-linked ***immunoassay*** was used to determine ***antibodies*** to HP in serum. Results: The overall prevalence of HP infection was 72%. In a multivariant discriminant analysis: age, country of origin and ethnic group were found to be the most closely associated variables for HP infection and the discriminant analysis succeeded in predicting correctly, in 62% of the population, whether they had or did not have HP infection. There was no significant difference with gender, occupation, educational level, blood group, smoking, gastrointestinal symptoms and use of medication. Conclusions: The prevalence of HP infection was higher than that in industrialized countries, but lower than in developing countries. The prevalence in a rural population was slightly higher than that of an urban population in Israel (65%). The country of origin and ethnic group influenced the prevalence of HP infection and not birth and growing up on the Kibbutz.

L11 ANSWER 24 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1994:392492 BIOSIS

DN PREV199497405492

TI A new method for detecting anti Helicobacter ***pylori***

antibodies: An analytical and clinical evaluation.

AU Plebani, M. (1); Basso, D.; Brigato, L.; Cassaro, M.; Farinati, F.; Di Mario, F.; Rugge, M.

CS (1) Ist. Med. Lab., c/o Lab. Centrale, Via Giustiniani 2, 35128 Padova Italy

SO Journal of Clinical Laboratory Analysis, (1994) Vol. 8, No. 4, pp. 219-222.

ISSN: 0887-8013.

DT Article

LA English

AB The diagnosis of Helicobacter ***pylori*** (Hp) infection is an important goal in clinical practice. In this paper we evaluated 1. the analytical reliability of a new second-generation antigen based enzyme ***immunoassay*** (Cobas Core Anti Helicobacter ***pylori*** EIA) in detecting anti-Hp IgG ***antibodies***, and 2. the behaviour of anti-HP IgG in patients with chronic atrophic and non-atrophic ***gastritis*** as compared to healthy controls. The findings from the

dilution curve, the values of intra and inter assay coefficients of variations (never above 10%) and of the recovery test (between 96 and 109%), confirm that the method is reliable. Serum IgG anti-Hp levels were found to be significantly higher in patients with histologically identified Hp infection, than in those negative at histology. Furthermore, the grade of histological positivity was correlated with serum IgG levels. However, we found a discrepancy between a low prevalence of Hp staining and a high prevalence of Hp seropositivity in patients with chronic atrophic or non-atrophic ***gastritis***, but not in controls. This suggests that IgG serum determination may be more useful than histology in determining a present or previous infection in patients with chronic atrophic or non-atrophic ***gastritis***.

L11 ANSWER 25 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1994:359852 BIOSIS

DN PREV199497372852

TI Comparison of PCR and other diagnostic techniques for detection of Helicobacter ***pylori*** infection in dyspeptic patients.

AU Weiss, Judith (1); Mecca, James; Da Silva, Elvira; Gassner, Dieter

CS (1) Dep. Infect. Dis., Roche Mol. Systems, Alameda, CA 94501 USA

SO Journal of Clinical Microbiology, (1994) Vol. 32, No. 7, pp. 1663-1668. ISSN: 0095-1137.

DT Article

LA English

AB A sensitive and specific PCR-based assay to detect the Helicobacter ***pylori*** 16S rRNA gene present in formalin-fixed paraffin-embedded ***gastric*** biopsy specimens has been developed. A total of 95 patients with dyspepsia were evaluated for the presence of chronic active ***gastritis*** and an infection with H. ***pylori*** through the use of diagnostic assays based on biopsy specimens and serology. The "gold standard" for the presence of the bacteria was direct detection in histological sections of biopsy specimens by Giemsa stain. The results obtained with the PCR assay performed on the biopsy specimens (94% sensitivity and 100% specificity) were equivalent to the detection of H. ***pylori*** immunoglobulin G ***antibodies*** by the commercially available second-generation Cobas Core anti-H. ***pylori*** immunoglobulin G enzyme ***immunoassay*** (94% sensitivity and 98% specificity) for the diagnosis of H. ***pylori*** infection. Urease testing and bacterial culture of the biopsy specimens were inferior (88 and 70% sensitivity and 96 and 98% specificity, respectively). A Western blot (immunoblot) analysis had slightly greater sensitivity (96%), although specificity was reduced to 93%. This research prototype PCR assay was shown to be highly reliable for the detection of infection with H. ***pylori*** and the presence of chronic active ***gastritis*** in the population studied.

L11 ANSWER 26 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1994:78004 BIOSIS

DN PREV199497091004

TI Evaluation of a new immunodiagnostic assay for Helicobacter ***pylori*** ***antibody*** detection: Correlation with histopathological and microbiological results.

AU Pronovost, Allan D. (1); Rose, Steven L.; Pawlak, Jan W.; Robin, Howard; Schneider, R.

CS (1) Quidel Corp., 10165 McKellar Court, San Diego, CA 92121 USA

SO Journal of Clinical Microbiology, (1994) Vol. 32, No. 1, pp. 46-50. ISSN: 0095-1137.

DT Article

LA English

AB Infection with Helicobacter ***pylori*** has been associated with the pathogenesis of chronic active ***gastritis*** and ***gastric*** and duodenal ulcer disease. Detection of immunoglobulin G ***antibodies*** to H. ***pylori*** offers a simple alternative to direct detection of the organism in biopsied tissue by culture or histopathological methods. A rapid flow-through membrane-based enzyme ***immunoassay*** for the detection of human immunoglobulin G ***antibodies*** to H. ***pylori*** has been developed and evaluated. Clinical evaluations were performed with 256 patient serum samples obtained from four clinical sites. Biopsy samples were obtained by endoscopic procedures at the same time as the serum samples, and were histopathologically and microbiologically categorized for the presence or absence of H. ***pylori*** . Sensitivity and specificity for this rapid enzyme ***immunoassay*** were 92 and 88%, respectively, compared directly with endoscopy results. After discordant results were resolved by a quantitative microwell enzyme-linked immunosorbent assay, the resulting sensitivity and specificity were 94 and gt 99%, respectively. These results indicate that this rapid enzyme ***immunoassay*** is a useful technique to determine H. ***pylori*** infection status and is a viable alternative to invasive endoscopic procedures.

L11 ANSWER 27 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1993:423557 BIOSIS

DN PREV199345071182

- TI Rapid test for detection of ***antibodies*** to Helicobacter
 pylori
- AU Anderson, G.; Alemohammad, M. M.; Foley, T. J.; Colletti, A.; Patel, A.; Dooley, C. P.
- CS Hycor Biomed. Inc., 7272 Chapman Ave., Garden Grove, CA 92641 USA
- SO Clinical Infectious Diseases, (1993) Vol. 16, No. SUPPL. 4, pp. S416-S417. Meeting Info.: First North American Congress on Anaerobic Bacteria and Anaerobic Infections Marina del Rey, California, USA July 24-26, 1992 ISSN: 1058-4838.

DT Article

LA English

L11 ANSWER 28 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1993:344308 BIOSIS

DN PREV199396041308

- TI Diagnosis of Helicobacter ***pylori*** infection by using Pyloriset EIA-G and EIA-A for detection of serum immunoglobulin G (IgG) and IgA ***antibodies***.
- AU Granberg, Christer (1); Mansikka, Antti; Lehtonen, Olli-Pekka; Kujari, Harry; Gronfors, Reijo; Nurmi, Heimo; Raihae, Ismo (1); Stahlberg, Marja-Riitta; Leino, Rauli
- CS (1) Orion Corp., Orion Diagnostica, SF-02101 Espoo Finland
- SO Journal of Clinical Microbiology, (1993) Vol. 31, No. 6, pp. 1450-1453. ISSN: 0095-1137.

DT Article

LA English

AB We evaluated the performance of new enzyme ***immunoassay*** (EIA)

kits (Pyloriset; Orion Corporation, Orion Diagnostica, Espoo, Finland) for the detection of immunoglobulin G (IgG) and IgA ***antibodies*** to Helicobacter ***pylori*** in serum. Serum samples from 195 patients with upper abdominal complaints were collected. Biopsy specimens of the ***gastric*** mucosae were taken for histological analysis and bacterial culture. The sensitivity, specificity, positive and negative predictive values, and efficacy of the Pyloriset EIA-G in detecting IgG ***antibodies*** to H. ***pylori*** were 92, 84, 88, 90, and 89%, respectively, when compared with those of the reference methods used. The corresponding data for detection of IgA ***antibodies*** were 80, 89, 89, 79, and 84%, respectively. The overall prevalence of defined H. ***pylori*** positivity was 54%. Moreover, the ***antibody*** tests showed a very good correlation with the biopsy findings. IgG ***antibodies*** were found in 93% of sera from patients with documented ***gastritis*** and H. ***pylori*** positivity, whereas only 4% of the sera from patients with documented ***gastritis*** and H. ***pylori*** -negative patients was positive. The results obtained for IgA ***antibodies*** were 81 and 6%, respectively. We conclude that the Pyloriset EIA-G, the test for IgG ***antibodies***, is a good and reliable test for the detection of ***antibodies*** to H. ***pylori*** and as an indication of H. ***pylori*** infection. The determination of IgA ***antibodies*** may be used as a test that complements the IgG ***antibody*** assay.

L11 ANSWER 29 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1992:524993 BIOSIS

DN BA94:133068

TI SERODIAGNOSIS OF HELICOBACTER- ***PYLORI*** INFECTIONS WITH AN ENZYME
IMMUNOASSAY USING THE CHROMATOGRAPHICALLY PURIFIED 120 KILODALTON
PROTFIN

AU GERSTENECKER B; ESCHWEILER B; VOEGELE H; KOCH H K; HELLERICH U; KIST M CS INST. MED. MICROBIOL., UNIV. FREIBURG, HERMANN-HERDER-STR. 11, 7800 FREIBURG, GER.

SO EUR J CLIN MICROBIOL INFECT DIS, (1992) 11 (7), 595-601. CODEN: EJCDEU. ISSN: 0934-9723.

FS BA; OLD

LA English

AB A membrane-associated 120 kDa protein on Helicobacter ***pylori*** with known species-specificity was isolated and used in an enzyme ***immunoassay*** (EIA) for the detection of Helicobacter ***pylori*** -specific IgG ***antibodies*** in patient sera. The EIA was compared with two other methods used for serodiagnosis of Helicobacter ***pylori*** infections: an EIA using sonicated whole Helicobacter ***pylori*** cell antigen and Western immunoblot. In a pospective study 127 unselected patients (76 patients with antrum ***gastritis***, 51 patients without ***gastritis***) who underwent gastroscopy were studied histologically and serologically. The EIA using the purified 120 kDa protein had the highest specificity (92%) compared with the EIA using a whole cell sonicate of a single Helicobacter ***pylori*** strain as antigen helicobacter ***pylori*** strain as antigen (60.7%) and the immunoblot (90.2%). The sensitivity was 96%, 100% and 92%, respectively. Sera of three control patients reacted strongly in all three methods, indicating possible Helicobacter ***pylori*** infection with negative histological finding. The EIA using the 120 kDa protein as antigen was shown to be a specific and sensitive technique for the serodiagnosis of

L11 ANSWER 30 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS AN 1992:524992 BIOSIS DN BA94:133067 TI SERODIAGNOSIS OF HELICOBACTER- ***PYLORI*** INFECTIONS BY DETECTION OF IMMUNOGLOBULIN G ***ANTIBODIES*** USING AN IMMUNOBLOT TECHNIQUE AND ENZYME ***IMMUNOASSAY*** AU FAULDE M; SCHROEDER JP; SOBE D CS ERNST-RODENWALDT-INSTITUT, FACHBEREICH II MED. MIKROBIOLOGIE, VIKTORIASTR. 11-13, 5400 KOBLENZ, GER. SO EUR J CLIN MICROBIOL INFECT DIS, (1992) 11 (7), 589-594. CODEN: EJCDEU. ISSN: 0934-9723. FS BA; OLD LA English AB A transferable solid phase enzyme ***immunoassay*** (TSP-EIA) and an immunoblot technique were evaluated for the detection of IgG ***antibodies*** against Helicobacter ***pylori***. Using the biopsy urease test as reference method, the sensitivity and specificity of the EIA were 96% and 100%, respectively. Immunoblot analysi was carried out by testing sera from patients with a positive urease test who suffered from type B ***gastritis***, ***gastric*** and duodenal ulcers, and a negative control group. The immunoblotted Helicobacter ***pylori*** proteins showed reproducible immunoreactive bands at molecular weights of 130, 93, 75 and 67 kDa. The molecular weight protein fractions of Helicobacter ***pylori*** of 180 kDa and higher were found to be of minor immunological significance. Proteins of less than 60 kDa exhibited wide serum-specific variations in reactivity after immunostaining. No correlation between specific immunoblot patterns and clinical signs induced by Helicobacter ***pylori*** infection was observed. L11 ANSWER 31 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS AN 1992:96603 BIOSIS DN BA93:53153 TI EVALUATION OF A COMMERCIALLY AVAILABLE SECOND-GENERATION IMMUNOGLOBULIN G ENZYME ***IMMUNOASSAY*** FOR DETECTION OF HELICOBACTER- ***PYLORI*** INFECTION. AU GOOSSENS H; GLUPCZYNSKI Y; BURETTE A; VAN DEN BORRE C; BUTZLER J-P CS WORLD HEALTH ORGANIZATION COLLABORATING CENTRE ENTERIC CAMPYLOBACTER, ST-PIETERS UNIVERSITY HOSPITAL, HOOGSTRAAT 322, B-1000 BRUSSELS, BELGIUM. SO J CLIN MICROBIOL, (1992) 30 (1), 176-180. CODEN: JCMIDW. ISSN: 0095-1137. FS BA; OLD LA English AB We evaluated a commercially available second-generation anti-H. ***pylori*** immunoglobulin G enzyme ***immunoassay*** (EIA) (Cobas Core Anti-Helicobacter ***pylori*** EIA; Roche S.A., Basel, Switzerland) for serodiagnosis of H. ***pylori*** infection. The results of the assay were assessed in relation to the results of bacterial culture, urease testing, and histological Giemsa stain of ***gastric*** biopsy specimens from 1,134 patients with a variety of symptoms relating to the upper gastrointestinal tract. H. ***pylori*** was detected in biopsy specimens from 660 (58.2%) patients: 6 had a normal mucosa, 123 had chronic ***gastritis*** only, and 531 were found to have chronic

active ***gastritis*** by histology; endoscopy showed duodenal and

gastric ulcers in 137 and 64 patients of the last two groups, respectively. The test was evaluated with different age and ethnic groups. The prevalence, sensitivity, specificity, and positive and negative predictive values were, respectively, (i) for Belgian patients between 18 and 40 years old, 34, 93, 95, 91, and 96%; (ii) for Belgian patients more than 40 years old, 53, 96, 91, 93, and 95%; and (iii) for Mediterranean patients more than 17 years old, 87, 94, 70, 95, and 64%. All sera showing discordant ***immunoassay*** results compared with the results of histology and culture of biopsy specimens, as well as those with borderline ***immunoassay*** results, were tested further by immunoblotting. Among the EIA results considered false negative, we demonstrated an absence of seroconversion in 14 of 19 patients tested by immunoblotting. Among the EIA results considered false positive, immunoblotting showed the presence of specific ***antibodies*** in 28 of 37 patients tested. Among the borderline results obtained in the first assay with 22 patients' sera, a second assay showed positive results in 10 patients (8 were positive by immunoblotting) and negative reactions in 10 patients (9 were negative by immunoblotting), whereas 2 remained borderline. These data indicate that sera showing borderline ***immunoassay*** results must be tested again. In conclusion, this commercially available second-generation EIA, which is easy and quick to perform, was found highly reliable for the serodiagnosis of H. ***pylori*** infection.

L11 ANSWER 32 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1991:402414 BIOSIS

DN BR41:64259

TI EVALUATION OF AN ENZYME ***IMMUNOASSAY*** MEASURING SERUM ***ANTIBODY*** TO HELICOBACTER- ***PYLORI*** VS. WESTERN BLOT AND HISTOCHEMICAL UREASE TESTING.

AU PASKELL S; HOUGHTON R; VIGOREN E; THRESHER K

CS BAINBRIDGE LAB. INC., BAINBRIDGE, IS., WA 98110.

SO 91ST GENERAL MEETING OF THE AMERICAN SOCIETY FOR MICROBIOLOGY, DALLAS, TEXAS, USA, MAY 5-9, 1991. ABSTR GEN MEET AM SOC MICROBIOL. (1991) 91 (0), 343.

CODEN: AGMME8.

DT Conference

FS BR; OLD

LA English

L11 ANSWER 33 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1991:388110 BIOSIS

DN BA92:65425

TI A NOVEL ENZYME ***IMMUNOASSAY*** FOR SERODIAGNOSIS OF HELICOBACTER-***PYLORI*** INFECTION.

AU SUGIYAMA T; IMAI K; YOSHIDA H; TAKAYAMA Y; YABANA T; YOKOTA K; OGUMA K; YACHI A

CS DEP. INTERNAL MEDICINE, SAPPORO MEDICAL COLLEGE, S-1, W-16, CHUO-KU, SAPPORO 060, JPN.

SO GASTROENTEROLOGY, (1991) 101 (1), 77-83. CODEN: GASTAB. ISSN: 0016-5085.

FS BA; OLD

LA English

AB Helicobacter ***pylori*** has recently been implicated as an etiologic agent of gastroduodenal disorders. Comparing the ***antibody*** to H.

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***pylori*** in the sera of patients with that of normal controls by
Western blot analysis, a unique ***antibody*** was detected in the
sera of patients, which reacted with the 25-kilodalton antigen of H.
 ***pylori*** . On the other hand, monoclonal ***antibody*** CP3
prepared in the authors' laboratory also recognized the 25-kilodalton
antigen of H. ***pylori*** . Whether the serum ***antibody***
the patient recognized the CP3 antigen purified by monoclonal
 ***antibody*** CP3 was then examined. Western blot analysis showed that
the patient's serum reacted strongly with the affinity-purified CP3
antigen. Using monoclonal ***antibody*** CP3, an enzyme-linked
immunosorbent assay to detect CP3 ***antibody*** in sera was
established. In patients with chronic ***gastritis*** and
 ***gastric*** ulcers, the titer of CP3 ***antibody*** was
significantly higher than in normal controls and correlated with the
histological grade of antral ***gastritis*** . The detection of CP3
 ***antibody*** in sera is useful in the diagnosis of chronic
 ***gastritis*** and ***gastric*** ulcer associated with H.
 ***pylori*** infection and also in evaluation of the grade
 ***gastritis*** .
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L11 ANSWER 34 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1990:355466 BIOSIS

DN BA90:52045

TI INOCULATION OF BARRIER-BORN PIGS WITH HELICOBACTER- ***PYLORI*** A USEFUL ANIMAL MODEL FOR ***GASTRITIS*** TYPE B.

AU ENGSTRAND L; GUSTAVSSON S; JORGENSEN A; SCHWAN A; SCHEYNIUS A

CS DEP. CLINICAL BACTERIOLOGY, UNIVERSITY HOSPITAL, UPPSALA, SWED.

SO INFECT IMMUN, (1990) 58 (6), 1763-1768. CODEN: INFIBR. ISSN: 0019-9567.

FS BA; OLD

LA English

AB At the age of 8 weeks, 15 barrier-born pigs, specific pathogen free, were inoculated intragastrically with suspension of 107 to 1010 CFU of Helicobacter ***pylori*** after treatment with omeprazole. The pigs were observed or up to 12 weeks, endoscopic biopsy specimens were taken, and serum samples were drawn. H. ***pylori*** was identified by routine culturing and by staining with an H. ***pylori*** -specific monoclonal ***antibody*** on cryostat sections of ***gastric*** biopsy specimens. In 11 of 15 inoculated pigs, H. ***pylori*** was detected throughout the observation period. In these infected pigs, there was an ***antibody*** response to H. ***pylori***, as determined in serum by an enzyme ***immunoassay*** . Furthermore, the development of superficial, focal ***gastritis*** with infiltrates of mononuclear class II antigen-expressing lymphocytes was observed immunohistologically. H. ***pylori*** was never detected and an ***antibody*** response to H. ***pylori*** was not observed in two control pigs. The development of ***gastritis*** and the systemic ***antibody*** response to H. ***pylori*** support the usefulness of this animal model for studies of H. ***pylori*** -related human diseases.

L11 ANSWER 35 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1990:218383 BIOSIS

DN BA89:115673

TI ISOTYPE EVOLUTION IN THE FOLLOW-UP STUDY OF PATIENTS WITH CAMPYLOBACTER-***PYLORI*** ASSOCIATED ***GASTRITIS***

AU GOBERT B; BENE M C; DE KORWIN J D; FAURE G CS LAB. D'IMMUNOL., CHRU DE NANCY, BP 184, F-54500 VANDOEUVRE-LES-NANCY, FR. SO GASTROENTEROL CLIN BIOL, (1989) 13 (11), 880-883. CODEN: GCBIDC. ISSN: 0399-8320. FS BA; OLD LA English AB Four sequential immuno-assays were performed from May to November 1988 to follow the levels of IgA, IgA and IgM to Campylobacter ***pylori*** in 16 infected patients with histologically proven ***gastritis***, among which 12 received appropriate therapy. Histopathological examination of antral biopsies, bacteriological cultures and urease tests were performed on each occasion when serum was tested for ***antibodies*** to C. ***pylori*** . The detection and quantitative assessment of the various isotypes to this bacterium proved valuable to appreciate the response to therapy with, in case of success, a steady decrease of ***antibodies*** levels concomitant with clinical improvement. L11 ANSWER 36 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS AN 1990:156704 BIOSIS DN BA89:84122 TI VALUE OF SEROLOGY ELISA AND IMMUNOBLOTTING FOR THE DIAGNOSIS OF CAMPYLOBACTER- ***PYLORI*** INFECTION. AU PENA A S; ENDTZ H P; OFFERHAUS G J A; HOOGENBOOM-VERDEGAAL A; VAN DUIJN W; DE VARGAS N; DEN HARTOG G; KREUNING J; VAN DER REYDEN J; ET AL CS DEP. GASTROENTEROL., LEIDEN UNIV. HOSP., P.O. BOX 9600, NL-2300 RC LEIDEN, SO DIGESTION, (1989 (1990)) 44 (3), 131-141. CODEN: DIGEBW. ISSN: 0012-2823. FS BA; OLD LA English AB Fifty-two unselected patients referred to for upper gastrointestinal endoscopy were evaluated in several ways to determine the presence of Campylobacter ***pylori*** . ***Antibodies*** against this microorganism were measured to assess the value of serology for the diagnosis of C. ***pylori*** infection. Five antral biopsy specimens were taken in each patient for culture and bacteriological determinations, histology [morphology and Warthin-Starry (WS) staining] and the urease test (2, 3 and 24 h). Serum ***antibodies*** against a sonicate of 6 strains of micoorganisms were assayed by enzyme-linked ***immunoassay*** (ELISA) and an immunoblotting technique. In 14 of the 52 patients the histology of the antrum was normal, 18 patients had chronic active ***gastritis*** and 20 had chronic ***gastritis*** without

strains of micoorganisms were assayed by enzyme-linked ***immunoassay (ELISA) and an immunoblotting technique. In 14 of the 52 patients the histology of the antrum was normal, 18 patients had chronic active ***gastritis*** and 20 had chronic ***gastritis*** without polymorphonuclear infiltration. In the group with normal histology, only 1 patient was positive for C. ***pylori*** with all methods, and 1 other subject was positive for IgG and 2 for IgA only with ELISA. In the group with chronic active ***gastritis***, 14 were positive with all methods, 1 was negative by WS only and another was negative for IgA according to ELISA, WS and ***antibodies***. Among the patients with chronic ***gastritis***, 7 were positive and 7 negative with all tests; in the other 6 patients the results obtained with the various tests were divergent. Four serological tests were studied and validated against culture, WS and urease test which were considered to be the reference methods. The serological tests showed high sensitivity and specificity for the detection of C. ***pylori*** -associated active chronic ***gastritis*** of the antrum, and can therefore serve as noninvasive

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L11 ANSWER 37 OF 184 BIOSIS COPYRIGHT 2001 BIOSIS
AN 1989:336534 BIOSIS
DN. BA88:39534
TI DIAGNOSTIC VALUE OF AN ***IMMUNOASSAY*** TO DETECT ANTI CAMPYLOBACTER-
    ***PYLORI*** ***ANTIBODIES*** IN NON-ULCER DYSPEPSIA.
AU LOFFELD R J L F; STOBBERINGH E; FRENDRIG J A; VAN SPREEUWEL J P; ARENDS J
CS DEP. INTERN. MED., UNIV. HOSPITAL MAASTRICHT, PO BOX 1918, 6201BX
   MAASTRICHT, NETH.
SO LANCET, (1989) 1 (8648), 1182-1185.
   CODEN: LANCAO. ISSN: 0140-6736.
FS BA; OLD
LA English
AB An enzyme-linked immunosorbent assay (ELISA) for detection of IgG
    ***antibodies*** against Campylobacter ***pylori*** was used to
   examine sera from 70 patients with non-ulcer dyspepsia. 48 patients had C
    ***pylori*** associated ***gastritis*** according to culture or
   histology, mean optical density (OD) of the ELISA was significantly higher
   than that for the 22 patients with normal antral mucosa and absence of C
    ***pylori*** . Positive and negative predictive values for
   campylobacter-associated ***gastritis*** were 100% above OD 2.10 and
  below OD 1.00, respectively. Serology might replace endoscopy in the
  diagnosis of campylobacter-associated ***gastritis***.
L11 ANSWER 38 OF 184 CAPLUS COPYRIGHT 2001 ACS
AN 2000:790743 CAPLUS
DN 133:319313
TI Method for assessing the risk of peptic ulcer, comprising the steps of
  determining quantitatively the concentrations of ***pepsinogen*** I
  (pgi) and ***gastrin*** -17 in a serum sample
IN Sipponen, Pentti; Harkonen, Matti; Suovaniemi, Osmo; Forsblom, Erik
PA Locus Genex Oy, Finland
SO PCT Int. Appl., 19 pp.
  CODEN: PIXXD2
DT Patent
LA English
FAN.CNT 1
  PATENT NO.
                  KIND DATE
                                    APPLICATION NO. DATE
PI WO 2000067035 A1 20001109 WO 2000-FI377 20000428
    W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR,
      CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU,
      ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU,
      LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE,
      SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA,
      ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
    RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
      DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
      CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
  FI 9900992
                A 20001031
                                FI 1999-992
                                              19990430
PRAI FI 1999-992 19990430
AB The present invention concerns a method for assessing the risk of peptic
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ulcer by detg. the presence and topog. phenotype of ***gastritis*** in

an individual, by detg. quant. the persinogen I and ***gastrin*** -17 concns. in a serum sample from the said individual, selecting a method-specific ref. value and cut-off value for resp. analyte, assessing the topog. and phenotype of ***gastritis*** based on a comparison of the ***pepsinogen*** I and ***gastrin*** -17 concns. so detd. with their resp. method-specific ref. and cut-off values, and correlating the so assessed ***gastritis*** phenotype with the risk for peptic ulcer. Preferably also Helicobacter ***antibodies*** are detd. in the sample.

RE.CNT 3

RE

- (1) Javier, P, European Journal of Gastroenterology & Hepatology 1999, V11,
- (2) Locus Genex Oy; WO 9615456 A1 1996 CAPLUS
- (3) Tseng-Shing, C; The American Journal of Gastroenterology 1994, V89(9), P1511
- L11 ANSWER 39 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 2000:275448 CAPLUS

DN 133:280240

- TI Serum reactivity to Helicobacter ***pylori*** antigens assessed by enzyme ***immunoassay*** and the immunoblot method in children
- AU Wojda, Urszula; Dzierzanowska, Danuta; Witkowska-Vogtt, Ewa; Celinska-Cedro, Danuta
- CS Department of Clinical Microbiology, Childrens Memorial Health Institute,
- SO Cent.-Eur. J. Immunol. (1999), 24(4), 257-264 CODEN: CJIMFW; ISSN: 1426-3912
- PB Polish Society for Immunology

DT Journal

LA English

AB H. ***pylori*** infection is the most common chronic bacterial disease in humans. It is the major cause of ***gastritis***, peptic ulcer, ***gastric*** carcinoma, and MALT lymphoma. The aim here was to assess the usefulness of 2 serol. methods, EIA and immunoblot, for detecting IgG ***antibodies*** to H. ***pylori*** in children. A total of 382 serum samples were included in the study. Group 1 contained 282 serum samples from children with dyspeptic symptoms of the proximal alimentary tract. Group 2 contained serum samples from 100 children without any pathol. symptoms in the alimentary tract. The results of serol. studies revealed that as many as 181/282 (64.2%) of symptomatic children showed the presence of specific IgG ***antibodies*** to H. ***pylori*** antigens. In contrast, in the group of asymptomatic children, only 27/100 (27%) showed the presence of anti-H. ***pylori*** IgG ***antibodies*** in the serum. Specific ***antibodies*** against protein CagA (116 kDa) were detected much more frequently in children with ***gastritis*** 40/45 (88.8%) and in children with duodenitis 3/4 (75%) than in those with normal ***gastric*** mucosa 8/15 (53%). Detection of anti-H. ***pylori*** ***antibodies*** by the use of serol. methods appeared helpful in diagnosis of infections induced by the bacteria.

RE.CNT 24

RE

- (2) Blaser, M; Cancer Res 1995, V55, P2111 CAPLUS
- (4) Covacci, A; Proc Natl Acad Sci USA 1993, V90, P5791 CAPLUS

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(10) Husson, M; J Clin Microbiol 1995, V33, P3300 CAPLUS
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(12) Lutton, D; J Med Microbiol 1995, V42, P386 CAPLUS

(15) Meijer, B; J Clin Microbiol 1997, V35, P292 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 40 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 2000:195874 CAPLUS

DN 133:55175

TI Detection of Helicobacter ***pylori*** using surface plasmon resonance

AU Nishimura, Tomoaki; Hifumi, Emi; Uda, Taizo

CS School of Biosciences, Hiroshima Prefectural University, Shoubara, 727-0023, Japan

SO Chem. Sens. (1999), 15(Suppl. A, Proceedings of the 28th Chemical Sensor Symposium, 1999), 148-150 CODEN: KAGSEU

PB Denki Kagakkai Kagaku Sensa Kenkyukai

DT Journal

LA Japanese

AB Helicobacter ***pylori*** (H. pyroli) causes chronic ***gastritis*** and ***gastric*** ulcer. It is important to detect the H. pyroli to administer of antibiotics for patients. We have established a unique monoclonal ***antibody*** which has a specificity against H. pyroli urease. Moreover it hugely suppresses the enzymic activity of the urease. In this study, H. pyroli and its sonicated samples were detected by using a differential SPR. The detection limit by the SPR was 2 x 107 cell/mL to the sonicated sample, which showed much higher detection limit than unsonicated H. pyroli by 100 fold.

L11 ANSWER 41 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 2000:19479 CAPLUS

DN 132:63138

TI Helicobacter ***pylori*** antigens applicable to diagnosing Helicobacter ***pylori*** infection

IN Kondo, Isamu, Hoshina, Sadayori, Miki, Keizaburo

PA Japan

SO Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PI JP 2000002706 A2 20000107 JP 1998-183354 19980615

AB New Helicobacter ***pylori*** antigen or its fragments are provided so that they are applied to diagnosing Helicobacter ***pylori*** infection without giving a pain to a patient and distinguishing ulcer and duodenal ulcer from ***gastritis***. These antigens are obtained from Helicobacter ***pylori*** present in ***gastric*** juice from ulcer patients or duodenal ulcer patients by the successive treatments with lysozyme and with N-acetylglucosaminidase, followed by electrophoresis. Their mol. wts. detd. by electrophoresis are apprx.68 kDa, apprx.78 kDa, apprx.132 kDa and apprx.140 kDa. The infection with Helicobacter ***pylori*** related to ulcer and duodenal ulcer, but not to ***gastritis***, was diagnosed by detecting the specific IgA ***antibodies*** present in serum with these antigens.

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DN 132:292571
TI Effect of cagA status on the sensitivity of enzyme ***immunoassay***
   in diagnosing Helicobacter ***pylori*** -infected children
AU Plebani, Mario; Guariso, Graziella; Fogar, Paola; Basso, Daniela; Gallo,
   Nicoletta; Zambon, Carlo Federico; Mozrzymas, Renata; Celadin, Marilena;
   Zacchello, Franco
CS Department of Laboratory Medicine, University Hospital of Padua, Italy
SO Helicobacter (1999), 4(4), 226-232
   CODEN: HELIFL; ISSN: 1083-4389
PB Blackwell Science, Inc.
DT Journal
LA English
AB The authors sought to evaluate in symptomatic children the influence of
   the Helicobacter ***pylori*** genotype on ***gastritis***,
   abdominal pain, and circulating anti-H. ***pylori*** IgG
    ***antibodies*** (anti-H. ***pylori*** IgG) or ***pepsinogen***
   A (PGA) and C (PGC). Also, they assessed anti-H. ***pylori*** IgG,
   PGA, and PGC patterns in a large cohort of asymptomatic children. The
   infection was found in 33 of 183 symptomatic children; among the 20 H.
    ***pylori*** -pos. children for which the H. ***pylori*** genotype
   was available, cagA was present or absent in equal percentages. H.
    ***pylori*** infection was assocd. with more severe ***gastritis***
   and higher serum levels of anti-H. ***pylori*** IgG and PGC but not
   with abdominal pain. In infected children, higher levels of anti-H.
    ***pylori*** IgG and the presence of abdominal pain were assocd. with
   infections caused by cagA-pos. strains. In the cohort of asymptomatic
   children, raised levels of anti-H. ***pylori*** IgG, PGA, and PGC were
   found in ~5% of the cases. Thus, infection with cagA-pos. H.
    ***pylori*** strains can be assocd. with increased frequency of reported
   abdominal pain and higher circulating levels of anti-H. ***pylori***
   IgG. The serol. assessment of H. ***pylori*** IgG using H.
    ***pylori*** antigens contg. significant amts. of cagA protein may,
   therefore, underestimate the true prevalence of infection.
RE.CNT 38
RE
(2) Atherton, J; J Biol Chem 1995, V270, P17771 CAPLUS
(4) Blaser, M; Cancer Res 1995, V55, P2111 CAPLUS
(5) Blaser, M; Cancer Res 1995, V55, P562 CAPLUS
(8) Cover, T; Mol Microbiol 1996, V20, P241 CAPLUS
(19) Hunter, F; Dig Dis Sci 1993, V38, P2081 CAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 43 OF 184 CAPLUS COPYRIGHT 2001 ACS
AN 1999:645846 CAPLUS
DN 132:219089
TI Anti-CagA reactivity in Helicobacter ***pylori*** -negative subjects: A
  comparison of three different methods
AU Fusconi, Marco; Vaira, Dino; Menegatti, Marcello; Farinelli, Silvia;
  Figura, Natale; Holton, John; Ricci, Chiara; Corinaldesi, Roberto;
  Miglioli, Mario
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CS Servizio di Patologia Medica II, Istituto di Clinica Medica I, University

of Bologna, Bologna, 40138, Italy

L11 ANSWER 42 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 1999:804109 CAPLUS

SO Dig. Dis. Sci. (1999), 44(8), 1691-1695 CODEN: DDSCDJ; ISSN: 0163-2116

PB Kluwer Academic/Plenum Publishers

DT Journal

LA English

AB Emerging evidence suggests that infection by CagA-pos. Helicobacter ***pylori*** strains is related to the development of more serious gastroduodenal diseases, thus conferring to the detn. of anti-CagA ***antibodies*** a relevant clin. significance in serol. screenings. The detection of anti-CagA positivity in sera neg. for anti-H. ***pylori*** ***antibodies*** raises the question of whether this apparently nonsense result is merely due to a false pos. reaction. To address this issue, we compared three different methods for the detection of anti-CagA ***antibodies*** . In all, 272 selected sera from patients with precisely defined H. ***pylori*** status (pos. or neg. concordance of five tests, ie, histol. by Giemsa in both antrum and corpus, rapid urease test, culture, [13C]urea breath test, IgG ELISA) were tested for anti-CagA reactivity by three different techniques (Western immunoblotting, ELISA, and recombinant immunoblotting assay). In order to assess the sensibility and specificity of each tests, we considered as "true" anti-CagA pos. sera those with two out of three pos. results. Sera from 70% of H. ***pylori*** -pos. patients and 10% from H. ***pylori*** -neg. patients turned out to be "true" positives for anti-CagA ***antibodies*** . The three methods showed similar excellent results, in terms of both sensitivity and specificity, always over 93%. It is confirmed that a proportion of patients with a neg. conventional serol. against H. ***pylori*** possess anti-CagA ***antibodies*** in their sera. In this paper we demonstrate that it can happen even in patients without any biol. signs of actual H. ***pylori*** infection. The possibility that this can be due to a false pos. lab. result is very likely ruled out by the accuracy of the three methods used. The clin. management of these patients needs further study on larger series.

RE.CNT 12

RE

- (1) Blaser, M; Lancet 1997, V349, P1020 MEDLINE
- (2) Censini, S; Proc Natl Acad Sci USA 1996, V93, P14648 CAPLUS
- (3) Covacci, A; Proc Natl Acad Sci USA 1993, V90, P5791 CAPLUS
- (4) Cover, T; J Clin Microbiol 1995, V33, P1496 MEDLINE
- (11) Towbin, H; Proc Natl Acad Sci USA 1979, V76, P4350 CAPLUS
- ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 44 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 1998:784058 CAPLUS

DN 130:137953

- TI Serological assessment of the early response to eradication therapy using an immunodominant outer membrane protein of Helicobacter ***pylori***
- AU Nishizono, Akira; Gotoh, Takayuki; Fujioka, Toshio; Murakami, Kazunari; Kubota, Toshihiro; Nasu, Masaru; Watanabe, Makoto; Mifune, Kumato
- CS Department of Infectious Diseases Control, Oita Medical University, Oita, 879-55, Japan
- SO Clin. Diagn. Lab. Immunol. (1998), 5(6), 856-861

CODEN: CDIMEN; ISSN: 1071-412X

PB American Society for Microbiology

DT Journal

LA English

AB Eradication of Helicobacter ***pylori*** infection cures ***gastritis*** and prevents recurrence of peptic ulcers. Endoscopy is usually used to evaluate the effectiveness of eradication therapy. We designed a new noninvasive assay system for the early evaluation of eradication of H. ***pylori*** infection in which a crude H. ***pylori*** outer membrane protein prepn. (HPOmp) is used as an antigen, and we detd. the sensitivity and specificity of the serol. assay system. Immunoblot anal. showed that anti-HPOmp ***antibodies*** reacted to a protein with a mol. mass of approx. 29 kDa. In those patients who responded to therapy, the anti-HPOmp IgG (IgG) titers measured by ELISA (ELISA) at 1 mo after the end of therapy were significantly lower than those before treatment (34.8% redn.; P < 0.001), and the posttreatment redn. in the ***antibody*** titer was significantly greater than that of the titer measured with a com. available anti-H. ***pylori*** IgG ELISA (34.8% vs. 16.1%; P < 0.001). When a 25% redn. of anti-HPOmp IgG titer at 1 mo after the end of treatment was taken as the cutoff value for H. ***pylori*** eradication, the sensitivity and specificity of our new assay were 75% (51 of 68 treatment responders) and 96% (22 of 23 nonresponders), resp. Our results indicate that the novel serol. test with HPOmp might be a clin. useful tool for assessment of eradication of H. ***pylori***.

RE.CNT 15

RE

- (2) Cullen, D; Lancet 1992, V340, P1161 MEDLINE
- (4) Evans, D; Infect Immun 1989, V57, P664 CAPLUS
- (5) Graham, D; Ann Intern Med 1992, V116, P705 MEDLINE
- (10) Nishizono, A; J Clin Microbiol 1993, V31, P1173 CAPLUS
- (15) Tomb, J, Science 1997, V388, P539 CAPLUS
- ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L11 ANSWER 45 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 1998:747526 CAPLUS

DN 130:51328

TI Reagent for detecting anti-Helicobacter ***pylori*** ***antibody***

IN Nishizono, Akira; Fujioka, Toshio; Mifune, Kumato; Watanabe, Makoto; Azumi, Junichi

PA Fujirebio, Inc., Japan

SO Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PI JP 10307142 A2 19981117 JP 1997-130552 19970506

AB The disclosed reagent comprises Helicobacter ***pylori*** -derived extracellular membrane 10.apprx.100 kDa mol. The 10.apprx.100 kDa Helicobacter antigen is useful for ***immunoassay*** in detecting Helicobacter ***pylori*** -specific ***antibody*** and for diagnosing Helicobacter infection, e.g. ***gastritis***

L11 ANSWER 46 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 1998:535180 CAPLUS

DN 129:215706

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TI Diagnostic agent for digestive tract diseases derived from Helicobacter
   infection
IN Yokota, Shinichi; Amano, Kenichi
PA Sumitomo Pharmaceuticals Co., Ltd., Japan
SO Jpn. Kokai Tokkyo Koho, 5 pp.
   CODEN: JKXXAF
DT Patent
LA Japanese
FAN.CNT 1
   PATENT NO.
                   KIND DATE
                                      APPLICATION NO. DATE
PI JP 10218900 A2 19980818
                                    JP 1997-32758 19970131
AB Disclosed is purified and immobilized Helicobacter ***pylori***
   lipopolysaccharide for diagnosing the seriousness of digestive tract
   diseases derived from Helicobacter infection. Serum samples derived from
   patients with chronic ***gastritis***, ***gastric*** ulcer and
    ***gastric*** cancer were tested with microplate contg. immobilized
   lipopolysaccharide for the presence of anti-Helicobacter
   lipopolysaccharide ***antibody*** .
L11 ANSWER 47 OF 184 CAPLUS COPYRIGHT 2001 ACS
AN 1998:197684 CAPLUS
DN 128:228250
TI Adhesins and adhesive proteins from Helicobacter ***pylori*** and
   their diagnostic and therapeutic uses
IN Ho, Bow
PA Cortecs International Limited, Australia; Chapman, Paul William; Ho, Bow
SO PCT Int. Appl., 26 pp.
   CODEN: PIXXD2
DT Patent
LA English
FAN.CNT 1
   PATENT NO.
                   KIND DATE
                                      APPLICATION NO. DATE
PI WO 9812562 A1 19980326
                                     WO 1997-GB2554 19970922
     W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
       DK, EE, ES, FI, GB, GE; GH, HU, ID, IL, IS, JP, KE, KG, KP, KR,
       KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ,
       PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG,
       US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
     RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR,
       GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA,
       GN, ML, MR, NE, SN, TD, TG
                  A1 19980414
                                    AU 1997-43116 19970922
   AU 9743116
   ZA 9708513
                  A 19990323
                                   ZA 1997-8513 19970922
PRAI GB 1996-19694 19960920
   GB 1996-22846 19961101
   WO 1997-GB2554 19970922
AB The present invention relates to novel methods of diagnosing Helicobacter
    ***pylori*** infection in a subject based on the immunol. interactions
   of H. ***pylori*** adhesive proteins and adhesin in particular. In
   addn., the invention relates to methods for distinguishing between
   different ***gastric*** disease states caused by H. ***pylori***
   including kits for use in such methods. Furthermore, the invention
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relates to the use of adhesin proteins in the prodn. of vaccines.

L11 ANSWER 48 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 1997:776267 CAPLUS

DN 128:58315

TI Cloning, detection, and, biological activity of Helicobacter

pylori ***gastric*** acid secretion inhibitory factor 1

IN Cave, David R.; Huang, Lili; Hoffman, James S.

PA St. Elizabeth's Hospital of Boston, USA

SO PCT Int. Appl., 48 pp. CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

PATENT NO. KIND DATE APPLICAT

APPLICATION NO. DATE

PI WO 9744464 A1 19971127 WO 1997-US8018 19970514 W: CA, JP

RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE PRAI US 1996-648986 19960517

AB The gene encoding acid inhibitory factor 1 (AIF-1) produced by Helicobacter ***pylori*** bacteria, was cloned and sequenced in its entirety and expressed in Escherichia coli. On the basis of a partial N-terminal amino acid sequence, a degenerate oligonucleotide was used to probe. an H. ***pylori*** gene library. An ***antibody*** to the N-terminal peptide was made, and an AIF-1 ELISA is described. AIF-1 is characterized as having a mol. mass of .apprx.90 kDa, an isoelec. point of 7.3, is inactivated by boiling temps. and the enzyme pronase. Its DNA sequence encodes a deduced amino acid sequence of 381 amino acids. Purified AIF-1 inhibits acid prodn. by ***gastric*** parietal cells.

L11 ANSWER 49 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 1996:731124 CAPLUS

DN 126:70729

- TI Immunomagnetic bead enrichment and PCR for detection of Helicobacter
 pylori in human stools
- AU Nilsson, Hans-Olof; Aleljung, Paer; Nilsson, Ingrid; Tyszkiewicz, Tadeusz; Wadstroem, Torkel
- CS Department of Medical Microbiology, University of Lund, Soelvegatan 23, S-223 62, Lund, Swed.
- SO J. Microbiol. Methods (1996), 27(1), 73-79 CODEN: JMIMDQ; ISSN: 0167-7012

PB Elsevier

DT Journal

LA English

AB An immunomagnetic bead-based polymerase chain reaction assay (IMS-PCR) was developed for the detection of Helicobacter ***pylori*** in exptl. inoculated human stools and human clin. stool samples. Magnetic beads coated with anti-H. ***pylori*** rabbit ***antibodies*** were used for enrichment and concn. of H. ***pylori*** from fecal samples. Taq polymerase inhibitors, found in human feces, are efficiently removed by the immunomagnetic sepn. (IMS) and subsequent washing of the magnetic beads. Conditions of the assay were developed and optimized with feces from a healthy, H. ***pylori*** seroneg., individual. Feces was inoculated with serial dilns. of either the spiral or the coccoid form of H. ***pylori***. These 2 morphol. forms could be detected at similar

concns. when inoculated in feces using an optimized IMS-PCR method. In 1 g of feces less than 2.5 times. 104 H. ***pylori*** cells were detected as measured with 2 sep. sets of PCR-primers, based on a urease A subunit gene sequence and a gene sequence encoding a 26-kDa surface protein of H. ***pylori*** Previously, no report has shown a sensitivity below 106 H. ***pylori*** in feces PCR. Preliminary anal. of stool samples from 17 patients with symptoms of ***gastritis*** and esophagitis by IMS-PCR showed a good correlation with EIA-anal. of H. ***pylori*** serum- ***antibodies*** from these patients. The results indicate that H. ***pylori*** cells are shed in feces of infected patients and that immunomagnetic bead PCR might be an appropriate method for clin. diagnosis and studies involving immunoprophylaxis, antibiotic treatment, as well as vaccine candidates.

L11 ANSWER 50 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 1996:453992 CAPLUS

DN 125:108455

- TI Novel ELISA kits for serum pepsinogens using polyclonal ***antibody***
 : Correlation with conventional RIA kits and clinical significance
- AU Matsukawa, Yoshihiro; Nishinarita, Susumu; Horie, Takashi; Matsumoto, Kyoici; Ishikawa, Masazumi; Hirooka, Tatsuo; Aoki, Takahito; Iwai, Chikara; Kurosaka, Hanzo
- CS School Medicine, Nihon University, Itabashi, 173, Japan
- SO Nihon Univ. J. Med. (1996), 38(3), 141-148 CODEN: NUMDAE; ISSN: 0546-0352

DT Journal

LA English

AB We established novel ELISA kits for estg. serum - *** pepsinogen *** Land ***pepsinogen*** II using polyclonal ***antibodies*** against pepsinogens. The correlations with results obtained with conventional kits for ***pepsinogen*** and the clin. significance of the novel kits were also evaluated. We measured the serum ***pepsinogen*** levels of endoscopically normal subjects and patients with gastroduodenal diseases employing both kits. The results for the serum ***pepsinogen*** levels measured with the ELISA kits correlated well with those measured with the conventional RIA kits (r=0.98 for both pepsinogens I and II). Using the ELISA kits, the serum ***pepsinogen*** II levels were found to be elevated in patients with ***gastric*** and duodenal ulcer as compared to those of endoscopically normal subjects (and 0.05). Concerning the serum ***pepsinogen*** I, patients with ***gastric*** polyp, chronic atrophic ***gastritis***, and intestinal metaplasia manifested lower levels than did normal subjects (0.01, and 0.01). The ratio of serum ***pepsinogen*** I to ***pepsinogen*** II were lower in patients with ***gastric*** cancer, ***gastric*** ulcer, ***gastric*** polyp, chronic atrophic ***gastritis***, and intestinal metaplasia as compared to those of in normal subjects (for ***gastric*** polyp, and 0.01 for the others). The ELISA kits were beneficial from the view point of radiohazard problems, and appeared to be useful tools for evaluating the condition of the gastroduodenal mucosa.

L11 ANSWER 51 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 1996:188384 CAPLUS

DN 125:55676

TI Helicobacter ***pylori*** infection. Part 2. Comparison of various serological tests

AU Haeckel, R.; Haeckel, Hella; Dirks, H.; Koessling, F.

CS Inst. Laboratoriumsmed., Zentralkrankenhaus St. Juergenstrasse, Bremen, D-28205, Germany

SO Laboratoriumsmedizin (1996), 20(2), 87-91 CODEN: LABOD3; ISSN: 0342-3026

DT Journal

LA German

AB IgG ***antibodies*** against H. ***pylori*** were detd. by the qual. Helisal Rapid Blood Test, and by 3 quant. tests (Enzygnost anti-Helicobacter ***pylori***, Helicobacter ***pylori***

Antibody test, and G.A.P. Test IgG). The estn. was divided in 2 groups (with and without acute ***gastritis*** caused by H.

pylori). The quant. ***antibody*** test gave the best results with 91% sensitivity and 96% specificity.

L11 ANSWER 52 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 1994:215316 CAPLUS

DN 120:215316

TI Rapid in vitro test and kit for Helicobacter ***pylori*** in saliva

IN Cripps, Allan W.; Stiel, Daniel; Witt, Campbell S.; Clancy, Robert L.

PA Auspharm International Ltd., Australia

SO Pat. Specif. (Aust.), 30 pp.

CODEN: ALXXAP

DT Patent

LA English

PI AU 644121

.FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

AU 1990-67676 19901203

B2 19931202

AU 9067676 A1 19910606 PRAI AU 1989-7718 19891204

AB A method for detecting contemporary infection by H. ***pylori*** in a mammal comprises contacting a mucous secretion from the mammal with an antigen component from H. ***pylori*** for a time and under conditions sufficient for an IgG ***antibody*** in the mucous secretion specific to the antigen component to form a complex and then subjecting the complex to a detecting means. The antigen component is immobilized onto a solid support (e.g., nitrocellulose membrane, glass, polymer). The antigen component of H. ***pylori*** comprises whole cell ext. and/or one or more isolated components thereof. The isolated component comprises a protein, a lipopolysaccharide, a polysaccharide, a lipid or any combination thereof. A kit is also disclosed. IgG ***antibodies*** to H. ***pylori*** were detd. in saliva samples by ELISA; the ***antibody*** levels were directly related to the level of H. ***pylori*** infection as indicated by the degree of ***gastric*** inflammation.

L11 ANSWER 53 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 1990:156555 CAPLUS

DN 112:156555

TI Detection of Campylobacter ***pylori*** urease ***antibodies***
and reagent therefor

IN Dent, Julie Claire

PA UK

SO PCT Int. Appl., 21 pp.

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DT Patent
LA English
FAN.CNT 1
  PATENT NO.
                  KIND DATE
                                     APPLICATION NO. DATE
PI WO 8909407
                 A1 19891005
                                    WO 1989-GB104 19890206
     W: AT, AU, BB, BG, BR, CH, DE, DK, FI, GB, HU, JP, KP, KR, LK, LU,
       MC, MG, MW, NL, NO, RO, SD, SE, SU, US
    RW: AT, BE, BJ, CF, CG, CH, CM, DE, FR, GA, GB, IT, LU, ML, MR, NL,
       SE, SN, TD, TG
   AU 8930547
                  A1 19891016
                                   AU 1989-30547 19890206
PRAI GB 1988-6831 19880323
   GB 1988-18531 19880804
   WO 1989-GB104 19890206
AB A reagent for use in the diagnosis of C. ***pylori*** infections
  comprises C. ***pylori*** urease attached to a solid surface, e.g. a
  microplate or latex beads. The reagent is used to detect
    ***antibodies*** to C. ***pylori*** urease in a serum sample. A
  test kit for use in diagnosis of C. ***pylori*** infections is
  provided. Thus, a std. ELISA protocol employing the above reagent was
  used to test for the presence of ***antibodies*** to C. ***pylori***
  urease in 202 patients receiving gastroscopy. There was a high
  correlation between the presence of proven ***gastritis*** and
   ***antibody*** to C. ***pylori*** urease. A very high
  discrimination was found between C. ***pylori*** neg. and pos.
  patients on microbiol. testing when the urease ELISA was used in testing
  serum. Eighteen sera from C. ***pylori*** -pos. patients which had
  been neg. on an ELISA using a sol. antigen gave high ***antibody***
  titers with the urease ELISA.
L11 ANSWER 54 OF 184 CAPLUS COPYRIGHT 2001 ACS
AN 1990:117176 CAPLUS
DN 112:117176
TI Antigenic compositions of Campylobacter ***pylori*** and methods for
  their production and diagnostic use
IN Blaser, Martin J.
PA USA
SO Eur. Pat. Appl., 23 pp.
  CODEN: EPXXDW
DT Patent
LA English
FAN.CNT 1
                  KIND DATE
                                    APPLICATION NO. DATE
  PATENT NO.
  ------
                                  EP 1989-400464 19890217
PI EP 329570
                 A2 19890823
  EP 329570
                A3 19910522
  EP 329570
                B1 19970502
    R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE
                                 US 1988-158003 19880218
  US 5459041
                 A 19951017
                                  CA 1989-591107 19890215
  CA 1339067
                 A1 19970729
                 A2 19900205
                                  JP 1989-39167 19890217
  JP 02035096
  JP 2738947
                B2 19980408
                                 AT 1989-400464 19890217
  AT 152524
                 E 19970515
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CODEN: PIXXD2

ES 2100850

T3 19970701

ES 1989-400464 19890217

PRAI US 1988-158003 19880218

AB ***Antibodies*** to C. ***pylori*** are detected using an antigenic compn. comprising fragments (esp. of the flagella) of 63, 57, 45, and/or 31 kilodaltons for diagnosis of ***gastritis*** or peptic ulcer disease. Levels of IgG, IgA, and IgM specific for C. ***pylori*** were detd. in blood serum samples, taken 8-581 days after ingestion of C. ***pylori***, by ELISA using polystyrene well-immobilized antigens of 5 strains of C. ***pylori***. Saroconversion in the IgA and IgG classes occurred between days 60 and 431 following challenge. For IgM there was a nearly 4-fold increase in optical d. between day 8 and 22 after challenge and a gradual decline afterwards. For diagnostic purposes, pos. threshold was 0.910, 0.470, and 2.6 optical d. units for IgG and IgA in blood sera and for IgG in urine, resp.

L11 ANSWER 55 OF 184 CAPLUS COPYRIGHT 2001 ACS

AN 1988:92717 CAPLUS

DN 108:92717

- TI Enzyme-linked immunosorbent assay of proton-potassium ***ATPase***, the parietal cell antigen
- AU Karlsson, F. A.; Burman, Pia; Loof, L.; Olsson, M.; Scheynius, Annika; Mardh, S.
- CS Dep. Med. Physiol. Chem., Univ. Uppsala, Uppsala, S-751-85, Swed.
- SO Clin. Exp. Immunol. (1987), 70(3), 604-10

CODEN: CEXIAL: ISSN: 0009-9104

DT Journal

LA English

AB Vesicular membranes, purified from porcine ***gastric*** mucosa and rich in H+, K+- ***ATPase***, were used to establish an enzyme-linked immunosorbent assay (ELISA) for detns. of parietal cell autoantibodies. Results obtained with the ELISA correlated well with std. immunofluorescence detns. of parietal cell ***antibodies*** based on frozen sections of rat stomach. The ELISA however was .apprx. 10-fold more sensitive than the immunofluorescence method and had high specificity. Intra- and interassay coeffs. of variation, detd. with a patient sera of av. positivity, were 5.5% and 18%, resp. The ELISA detected ***antibody*** binding in 23 of 26 sera from patients with known autoimmune atrophic ***gastritis***, in 5 of 25 sera with autoimmune thyroiditis, in 5 of 20 sera from patients with Graves' disease, in 3 of 20 sera from patients with atoxic nodular goiter, in 6 of 20 sera of patients with primary biliary cirrhosis, in 2 of 20 sera of patients with active duodenal ulcer, in 2 of 20 sera with detectable antinuclear ***antibodies***, and in 1 of 20 sera with detectable rheumatoid factor. Data detd. by an ELISA based on a ***gastric*** vesicular membrane prepn. of human origin correlated well to those obtained by the std. ELISA based on porcine membrane material. The assay should be well suited for routine detns. of parietal cell ***antibodies*** in investigations of autoimmune ***gastritis*** and multiple organ autoimmune endocrinopathies.

- L11 ANSWER 56 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 2000332591 EMBASE
- TI Analysis of Helicobacter ***pylori*** binding site on HEp-2 cells and three cell lines from human ***gastric*** carcinoma.
- AU Nishihara K.; Nozawa Y.; Nomura S.; Kitazato K.; Miyake H.
- CS K. Nishihara, Pharmacology Research Lab., Tokushima Research Center, Taiho

Pharmaceutical Co. Ltd., 224-2, Ebisuno, Hiraishi, Tokushima 771-0132, Japan SO Fundamental and Clinical Pharmacology, (1999) 13/5 (555-561). Refs: 36 ISSN: 0767-3981 CODEN: FCPHEZ CY France DT Journal; Article FS 004 Microbiology 016 Cancer 048 Gastroenterology LA English SL English AB Helicobacter ***pylori*** (H. ***pylori***) is a pathogen responsible for chronic ***gastritis*** and peptic ulcer diseases. It colonises the ***gastric*** mucus layer and adheres to the ***gastric*** epithelial cell surface. As this adherence is the first step of infection, it is important to study the adherence mechanism. The aim of this study was to analyse the specific binding assay of H. ***pylori*** to HEp-2 cells and three ***gastric*** phenotype cell lines, AGS, MKN-45 and AZ-521. H. ***pylori*** NCTC 11637 grown on agar plates was harvested and used in experiments. H. ***pylori*** was inoculated to pre-cultured cell monolayers. Adhered bacteria were labelled with an anti-H. ***pylori*** ***antibody*** and an FITC-conjugated secondary ***antibody*** and quantified by using a fluorescent plate reader. Microbial adherence to HEp-2 cells increased with incubation time and incubated concentration of H. ***pylori*** . No further increase was obtained with four or more hours of incubation or with a concentration of 4 x 107 bacteria/well or more. Scatchard analysis revealed a linear plot and the Bmax value was 88.3. Similar adherence patterns were obtained when AGS, AZ-521 and MKN-45 cells were used for adherence assays, but they had a lower binding affinity than HEp-2 cells and AZ-521. MKN-45 cells had less receptors than HEp-2 and AGS cells. In conclusion, H. ***pylori*** adhered to the cell surface could be quantified by this assay method. H. ***pylori*** adhesion to cell surfaces has a single population of binding site and one type of binding site on HEp-2, AGS, AZ-521 and MKN-45 cells. (C) 1999 Editions scientifiques et medicales Elsevier SAS. L11 ANSWER 57 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 2000301502 EMBASE TI The clinical role of stool test (HpSA) in noninvasive diagnosis of Helicobacter ***pylori*** infection. AU Vaira D.; Ricci C.; Acciardi C.; Gatta L.; Berardi S.; Miglioli M. CS Dr. D. Vaira, 1st Medical Clinic, University of Bologna, Nuove Patologie Policlin. S. Orsola, Bologna, Italy, vairadin@med.unibo.it SO Turkish Journal of Gastroenterology, (2000) 11/2 (97-102). Refs: 30 ISSN: 1300-4948 CODEN: TJGAF3 CY Turkey DT Journal: Article FS 004 Microbiology 006 Internal Medicine

Immunology, Serology and Transplantation

026 048

LA English SL English

Gastroenterology

AB Helicobacter ***pylori*** (H. ***pylori***) causes a chronic ***gastric*** infection, which is usually life-long and many epidemiological studies have shown that this is probably one of the most common bacterial infections throughout the world, involving 50% of the population in developed countries and up to 80-90% of the population in developing regions. It is therefore clean that nowadays the diagnosis of H. ***pylori*** infection today represents, at the very least, a key step in the management of many of the patients referred to the gastroenterologist. Due to the widerange and relevance of pathologies possibly related to infection (including malignancies), it also harbours the potential to become a major health problem. Up to now, there were only two widely available non-invasive methods: 1) 13C or 14C labelled urea breath test and 2) serology (which is based on the detection of a specific anti-H. ***pylori*** immune response, mostly by IgG ***antibodies*** , in patient's serum). Over the last few years H. ***pylori*** has been detected in the culture of stool samples but viable organisms are present only in a small percentage of cases. Despite the difficulties encountered in culture from stool samples, the fact that the organism was present at all raised the possibility of developing a new non-invasive diagnostic test based on the detection of bacterial antigen in stool. Over the last two years an enzymatic ***immunoassay*** (EIA), which detects the presence of H. ***pylori*** antigen in stool specimen has become available (HpSA(TM)- H. ***pylori*** Stool Antigen Meridian Diagnostics Inc., Cincinnati USA) and begun to be used in clinical practice to evaluate its performance compared to that of other currently available diagnostic tests. The HpSA test has recently received approval from the United States Food and Drugs Administration (FDA) for two indications for use: 1) diagnosis of H. ***pylori*** infection in adult symptomatic patients and 2) monitoring response and post-therapy in adult patients. The test utilises polyclonal anti-H. ***pylori*** capture ***antibody*** absorbed in microwells. It is clear that such a test, which detects bacterial antigen in an actual ongoing infection, is theoretically useful not only for screening, but also as an early predictor of successful treatment. This review will briefly consider the currently available evidence supporting a possible role for this non-invasive diagnostic test.

L11 ANSWER 58 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 2000258461 EMBASE

- TI Noninvasive tests as a substitute for histology in the diagnosis of Helicobacter ***pylori*** infection.
- AU Hahn M.; Fennerty M.B.; Corless C.L.; Magaret N.; Lieberman D.A.; Faigel D.O.
- CS Dr. D.O. Faigel, Division of Gastroenterology, Portland VA Medical Center (P3GI), 3710 SW US Veterans Hospital Rd., Portland, OR 97201, United States
- SO Gastrointestinal Endoscopy, (2000) 52/1 (20-26). Refs: 25

ISSN: 0016-5107 CODEN: GAENBQ

CY United States

DT Journal; Article

FS 004 Microbiology

- Biophysics, Bioengineering and Medical Instrumentation 027
- Health Policy, Economics and Management 036
- 048 Gastroenterology

LA English

SL English

AB Background: Rapid urease tests for Helicobacter ***pylori*** have a sensitivity of 80% to 90%. Therefore histologic examination of ***gastric*** biopsies is recommended as a 'backup' diagnostic test in rapid urease test- negative patients. However, noninvasive tests (urea breath test, serology, whole blood ***antibody*** tests) may provide a more rapid diagnosis and be less expensive but offer similar accuracy. Methods: Sixty-seven patients (no prior treatment for H ***pylori***, no proton pump inhibitors, antibiotics, or bismuth within 4 weeks) undergoing endoscopy for evaluation of dyspepsia symptoms and testing rapid urease test-negative by antral biopsy were enrolled. All had the following tests: ***gastric*** biopsies (2 antral, 1 fundus, H and E and Alcian Yellow stain) examined for ***gastritis*** and H ***pylori***; 13C-UBT; capillary blood for whole blood rapid ***antibody*** tests: FlexSure HP, QuickVue, AccuStat, and Stat-Simple ***Pylori***; serum for FlexSure HP; HM-CAP enzyme-linked ***immunoassay*** . H ***pylori*** infection was diagnosed (reference standard) if chronic ***gastritis*** was present on histology and at least 2 of the 3 following tests were positive: urea breath test, H ***pylori*** organisms unequivocally demonstrated in biopsies on special stain, and/or enzyme-linked ***immunoassay*** . The test and treatment costs per patient were calculated. Results: Of 67 patients with a negative rapid urease test, 4 were positive for H ***pylori*** . None had active peptic ulcer disease. Histology only identified I patient with organisms visible on special stain. Using chronic active ***gastritis*** (neutrophilic and mononuclear infiltrate) as a diagnostic criterion for H ***pylori***, 6 patients would have been judged positive. However, only 2 of these were truly positive by the reference standard (positive predictive value 33%). Negative predictive value for presence of organisms and chronic active ***gastritis*** was 95% and 97%, respectively. All of the noninvasive tests identified all 4 truly positive patients correctly. Urea breath test and FlexSure whole blood assay yielded a substantial number of false-positive results (positive predictive value 31% and 36%, respectively); positive predictive value for the other tests ranged from 50% to 80%. All tests except histology had a negative predictive value of 100%. Histology was the most costly test (p < 0.001compared with all other tests), followed by urea breath test and HM-CAP serology (p < 0.001 compared with all rapid ***antibody*** tests). Conclusions: Whole blood or serum ***antibody*** testing is a rapid, accurate, and cost-effective means for establishing H ***pylori*** status in rapid urease test- negative patients. Whole blood or serology rapid ***antibody*** testing should substitute for histology when the patient has not been previously treated for H ***pylori***.

- L11 ANSWER 59 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 2000206472 EMBASE
- TI Telomerase expression, Hp infection and ***gastric*** mucosal carcinogenesis.
- CS Dr. X.X. He, Department of Gastroenterology, First Affiliated Hospital, Sun Yat-sen Univ. of Med. Sciences, 58 Zhongshanerlu, Guangzhou 510080, Guangdong Province. hexingxiang@263.net
- SO World Chinese Journal of Digestology, (2000) 8/5 (505-508).

Refs: 11

ISSN: 1009-3079 CODEN: SHXZF2

CY China

DT Journal; Article

FS 004 Microbiology

005 General Pathology and Pathological Anatomy

016 Cancer

048 Gastroenterology

LA Chinese

SL English; Chinese

AB AIM: To analyze telomerase activity and whether it is implicated in Hp infection as well as the relationship among telomerase expression, Hp infection and ***gastric*** mucosal carcinogenesis. METHODS: Telomerase activity was detected by TRAP in normal ***gastric*** mucosa, precancerous lesions and ***gastric*** carcinoma. Serum Hp-CagA-IgG ***antibody*** was determined by EIA in Hp infected patients. The relationship between telomerase activity and Hp-CagA- IgG ***antibody*** was studied by match method using 22 pairs of Hp positive patients including corresponding non-neoplastic ***gastric*** mucosa of ***gastric*** carcinoma and chronic superficial ***gastritis*** (CSG) mucosa. RESULTS: All normal ***gastric*** mucosa and CSG were telomerase negative. The positive rates of telomerase activity of grade 0 (n = 20), 1 (n = 40) and 2 (n = 8) intestinal metaplasia (IM) are 0%, 25% and 38%, respectively. In 68 chronic atrophic ***gastritis*** (CAG). 16 of 18 ***gastric*** carcinoma showed telomerase activity, the positive rate was the highest 89% in all the biopsy specimens. Thirty-nine of 45 tumors had telomerase activity (89%). The positive rates of telomerase activity of IM grade 0 (n = 15), 1 (n = 22) and 2 (n = 8) were 0%, 32% and 100%, respectively in corresponding nontumorous tissues. The incidence of IM grade 2 or tumor specimens was significantly higher than that in IM grade 0 or 1 (P<0.01). The Hp positive rates at normal ***gastric*** (n = 10), CSG (n = 46), CAG IM grade 0 (n = 20), 1 (n = 46)40) and 2 (n = 8) were 0%, 52%, 60%, 70% and 75%, respectively. Hp infection increased as the grade of IM advanced, in parallel with telomerase expression in the CAG. Hp-CagA-IgG ***antibody*** in CSG patients was significantly lower than that in patients with ***gastric*** carcinoma (P<0.01). All of 22 Hp positive ***gastric*** carcinoma were CagA+ strains (100.%). Twelve (55%) of the 22 corresponding nontumorous ***gastric*** mucosa had positive telomerase activity. In contrast, only 8 of 22 Hp positive CSG were infected CagA+ strains (36%); and all of the 22 *** gastric*** mucosa showed negative telomerase. CONCLUSION: The normal and CSG mucosa express no telomerase activity. Telomerase activity expression increases as the grade of IM advanced in CAG. Telomerase activity of ***gastric*** carcinoma is the highest (>88%) in all the ***gastric*** mucosa. The degree of Hp infection increases in parallel with telomerase positively in corresponding non-cancerous mucosa of ***gastric*** cancer. Infected Hp is usually CagA+ strain in cancer, whereas infected Hp is usually CagA- strain in CSG. Thus, telomerase activity may serve as a powerful tool for an early ***gastric*** carcinoma diagnosis. CagA+ Hp infection may be a strong trigger for telomerase reactivation.

L11 ANSWER 60 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 2000144389 EMBASE

TI Diagnosis of Helicobacter ***pylori*** infection in patients with atrophic ***gastritis*** : Comparison of histology, 13C-urea breath test, and serology.

AU Kokkola A.; Rautelin H.; Puolakkainen P.; Sipponen P.; Farkkila M.; Haapiainen R.; Kosunen T.U.

CS Dr. P. Puolakkainen, Second Dept. of Surgery, Helsinki University, Central Hospital, Haartmaninkatu 4, FIN-00290 Helsinki, Finland

SO Scandinavian Journal of Gastroenterology, (2000) 35/2 (138-141).

Refs: 27

ISSN: 0036-5521 CODEN: SJGRA4

CY Norway

DT Journal; Article

FS 004 Microbiology

005 General Pathology and Pathological Anatomy

016 Cancer

026 Immunology, Serology and Transplantation

048 Gastroenterology

LA English

SL English

AB Background: Atrophic ***gastritis***, a risk factor for ***gastric*** cancer, is a late consequence of Helicobacter ***pylori*** infection in approximately one-third of the infected patients. It has been suggested that ***gastric*** cancer would develop less frequently if H. ***pylori*** were eradicated. However, the prevalence of H. ***pylori*** infection may be underestimated in patients with atrophic ***gastritis*** and intestinal metaplasia if only biopsy-based diagnostic methods are used. Methods: We compared histology, 13C-urea breath test (13C-UBT), and serology in H. ***pylori*** diagnostics in 50 male patients with atrophic corpus ***gastritis*** . Results: H. ***pylori*** was detected in 15 (30%)patients by histology and in 14 (28%) by 13C-UBT, whereas increased serum ***antibody*** levels indicating H. ***pylori*** infection were found in 41 (82%) patients (P < 0.0001 between serology and both histology and 13C-UBT). H. ***pylori*** infection was associated with atrophic corpus ***gastritis*** in 84% of the present patients (in one patient with normal ***antibody*** titres H. ***pylori*** was defined histologically). Conclusions: H. ***pylori*** infection would have been missed in most patients with atrophic ***gastritis*** without the analysis of H. ***pylori*** ***antibodies***. Therefore, in patients with atrophic ***gastritis***, the use of serology is encouraged in diagnosing H. ***pylori*** infection.

L11 ANSWER 61 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 1999423461 EMBASE

TI Use of serology, the urease test and histology in diagnosis of Helicobacter ***pylori*** infection in symptomatic and asymptomatic Indians.

AU Kang G.; Rajan D.P.; Patra S.; Chacko A.; Mathan M.M.

CS Dr. G. Kang, Department Gastrointestinal Sciences, Christian Medical College Hospital, Vellore 632004, India

SO Indian Journal of Medical Research, (1999) 110/SEP. (86-90). Refs: 17

ISSN: 0971-5916 CODEN: IMIREV

CY India

DT Journal; Article

FS 004 Microbiology

048 Gastroenterology

LA English

SL English

AB Age-specific prevalence of IgA and IgG ***antibodies*** in 714 subjects without gastrointestinal complaints aged 6 months to 90 yr was measured by an enzyme linked ***immunoassay*** using an acid-glycine extract of H. ***pylori*** as the antigen. The urease test and . histology were used for the diagnosis of H. ***pylori*** infection in 83 subjects with a clinical diagnosis of dyspepsia, and these results were compared with measurement of IgG, IgA and IgM ***antibodies*** . The age specific prevalence of IgG and IgA ***antibodies*** respectively was 57 and 43 per cent for subjects aged 6 months to 4 yr and showed an increase with age to a maximum of 90 per cent for IgG in subjects > 60 yr of age and to 87 per cent for IgA in subjects between 51 and 60 yr. In symptomatic patients, there was a high degree of correlation between severity of H. ***pylori*** infection on histopathological examination and IgG (P < 0.02) levels. The use of IgG and IgA estimation could have identified H. ***pylori*** infection without endoscopy in 50 of the 83 patients. Serology for IgG and IgA ***antibodies*** against H. ***pylori*** may play a major role in decreasing the need for endoscopy, but cut-off values must be determined for each assay based on the prevalence of ***antibodies*** in the population.

L11 ANSWER 62 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 1999383622 EMBASE

TI Monitoring of Helicobacter ***pylori*** eradication by anti-H. ***pylori*** determination.

AU Antoljak N.; Vulkadinovic M.V.; Zubcic A.; Topic E.

CS Dr. N. Antoljak, Clinical Institute of Chemistry, Sestre Milosrdnice Univ. Hospital, Vinogradska c. 29, HR-10000 Zagreb, Croatia

SO Acta Clinica Croatica, (1999) 38/3 (203-207).

Refs: 10

ISSN: 0353-9466 CODEN: ACLCED

CY Croatia

DT Journal; Article

FS 004 Microbiology

005 General Pathology and Pathological Anatomy

027 Biophysics, Bioengineering and Medical Instrumentation

048 Gastroenterology

LA English

SL English; Serbo-Croatian

AB Helicobacter ***pylori*** (H. ***pylori***) infection results in ***gastric*** mucosa inflammation called chronic superficial ***gastritis***, developing peptic ulceration in some patients. The two major categories of diagnostic tests are non-invasive tests and invasive methods using endoscopy. The aim of the study was to monitor the efficacy of anti-H. ***pylori*** by the non-invasive quantitative serologic method. The ELISA Pyloriset EIA-G (Orion Diagnostica, Finland) with a cut-off titer of 300 mg/L was used to measure the concentration of specific anti-H. ***pylori*** IgG in sera of 34 patients with positive dyspeptic illness history before, and then two, four and six months after the treatment. A titer decline by .gtoreq.40% of the ValUe measured before therapy is named seroconversion. The mean percentage of titer decline was greatest two months after the treatment (49%; p<0.001). A statistically significant decrease persisted after 4 and 6 months (66% and 78% of the pretherapeutic titer, respectively; p<0.005). After 6 month monitoring, 94% of patients were found to be successfully seroconverted. According to

some authors, a 4-month titer monitoring is needed to ascertain H.

pylori eradication and to confirm seroconversion. Unlike these studies, our results showed that 73.4% of patients had a significant titer decline after 4 months, while 94% of patients were seroconverted after 6 months. So, the ***antibody*** titer decrease recorded 6 months following antimicrobial treatment could be an indicator of successful eradication in almost all treated patients.

L11 ANSWER 63 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 1999328131 EMBASE

TI Helicobacter ***antibodies*** in Finnish centenarians.

AU Rehnberg-Laiho L.; Louhija J.; Rautelin H.; Jusufovic J.; Tilvis R.; Miettinen A.; Kosunen T.U.

CS Dr. T.U. Kosunen, Haartman Institute, Dept. of Bacteriology and Immunology, University of Helsinki, Haartmaninkatu 3, 00014 Helsinki, Finland. timo.kosunen@helsinki.fi

SO Journals of Gerontology - Series A Biological Sciences and Medical Sciences, (1999) 54/8 (M400-M403).

Refs: 25

ISSN: 1079-5006 CODEN: JGASFW

CY United States

DT Journal; Article

FS 020 Gerontology and Geriatrics

048 Gastroenterology

LA English

SL English

AB Background. The prevalence of helicobacter ***antibodies*** increases with age and, in many developed countries, is highest in people born before 1940. Data on very old subjects are, however, limited. In this study we wanted to determine whether the age-related increase in the seroprevalence of H. ***pylori*** infection continues even in the oldest age group alive in Finland, the centenarians. Methods. Sera from 173 subjects (93% of all centenarians alive in Finland in 1991) were available for the present study. IgG and IgA ***antibodies*** against H. ***pylori*** were determined by an in-house enzyme ***immunoassay*** To estimate the influence of atrophic ***gastritis*** on the prevalence of helicobacter ***antibodies***, serum ***pepsinogen*** I (PG I) concentrations and parietal cell ***antibodies*** (PCAs) were measured by an enzyme ***immunoassay*** and indirect immunofluorescence, respectively. Results. The prevalence of helicobacter ***antibodies*** in Finnish centenarians was 66%. Low PG I values (<28 .mu.g/l) were found in 36% and positive PCAs in 16% of the subjects studied. The prevalence of PCAs was especially high (50%) in H. ***pylori*** -negative subjects with low PG I values, suggesting severe ***gastric*** atrophy. Conclusions. The age-related increase in H. ***pylori*** seroprevalence did not continue in the oldest age group alive in Finland. This may be explained partly by a relatively high frequency of atrophic ***gastritis*** (as suggested by low PG I values) in H. ***pylori*** -negative centenarians, but other factors such as selective H. ***pylori*** -related mortality - may also have contributed to the fairly low seroprevalence (66%) observed.

L11 ANSWER 64 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 1999061826 EMBASE

TI Rapid immunochromatographic assay for detection of Helicobacter

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***pylori*** ***antibodies*** .
AU Pavlitou K.; Pastore F.; Moldovanidou K.; Gioula G.; Routsinas Ch.;
   Polidorou F.; Malaka E.
CS K. Pavlitou, Microbiol. 'Agios Demetrios' Dept., General Hospital,
   Thessaloniki, Greece
SO Acta Microbiologica Hellenica, (1998) 43/3 (267-270).
   Refs: 12
   ISSN: 0438-9573 CODEN: AMBHAA
CY Greece
DT Journal; Conference Article
FS 004
          Microbiology
   026
         Immunology, Serology and Transplantation
         Gastroenterology
   048
LA Greek
SL English; Greek
AB The aim of the study was to evaluate the accuracy of a rapid
   immunochromatographic assay Helicobacter ***pylori*** EASY-card in the
   qualitative determination of Helicobacter ***pylori*** (HP)
    ***antibodies*** . A total of 311 patients undergoing endoscopy was
   studied. Diagnosis of lip infection was established if rapid urease test
   and smear's stain (haematoxylineosin, Giemsa) were positive. Anti-HP
    ***antibodies*** were detected in the serum of patients using
   Helicobacter ***pylori*** EASY-card and enzyme ***immunoassay***
   (ELISA). The sensitivity and specificity of EASY-card and ELISA tests were
   64%, 69% and 89%, 79% respectively. The above results indicate that
   Helicobacter ***pylori*** EASy-card is a rapid and useful test;
   however, it should be used only as preliminary, alternative, non-invasive
   procedure in patients with suspected HP infection.
L11 ANSWER 65 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.
AN 1998419516 EMBASE
TI Positive association between Helicobacter ***pylori*** infection and
   food allergy in children.
AU Corrado G.; Luzzi I.; Lucarelli S.; Frediani T.; Pacchiarotti C.;
   Cavaliere M.; Rea P.; Cardi E.
CS Dr. G. Corrado, Paediatric Gastroenterology Unit, Paediatric Clinic
   Institute, La Sapienza University, Viale Regina Elena 324, I-00161 Rome,
SO Scandinavian Journal of Gastroenterology, (1998) 33/11 (1135-1139).
   Refs: 48
   ISSN: 0036-5521 CODEN: SJGRA4
CY Norway
DT Journal; Article
FS 004 Microbiology
         Pediatrics and Pediatric Surgery
  007
         Immunology, Serology and Transplantation
  048
         Gastroenterology
LA English
SL English
AB Background: In children Helicobacter ***pylori*** has been involved as
   a pathogenetic factor in ***gastritis*** and duodenal ulcer and as a
  cofactor in protein-losing enteropathy, chronic diarrhoea, short stature,
  and ***gastritis*** lymphoproliferative disease. A subset of an H.
    ***pylori*** strain possesses an antigen, CagA, as a virulence factor.
  In the present study we determined anti-H. ***pylori*** IgG and
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anti-CagA IgG titres in children with food allergy. Methods: Ninety paediatric patients were studied: 30 with food allergy, 30 with atopic asthma, and 30 with inflammatory bowel disease. Anti-H. ***pylori*** IgG and anti-CagA IgG were determined in all children by means of a commercial enzyme ***immunoassay*** (ELISA). Results: The anti-H. ***pylori*** IgG titre was significantly higher in allergic patients than in the other two groups. The anti-CagA IgG titre did not differ significantly between the patients. Conclusions: These findings show a positive association between H. ***pylori*** infection and food allergy in children. We hypothesize that virulence factors other than CagA may be involved in the pathogenesis of H. ***pylori*** infection in paediatric patients with food allergy.

L11 ANSWER 66 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 1998383390 EMBASE

TI Comparison of salivary and serum enzyme ***immunoassays*** for the diagnosis of Helicobacter ***pylori*** infection.

AU Embil J.M.; Choudhri S.H.; Smart G.; Aldor T.; Pettigrew N.M.; Grahame G.R.; Dawood M.R.; Bernstein C.N.

CS C.N. Bernstein, Division of Gastroenterology, GB443 Health Sciences Center, 820 Sherbrok Street, Winnipeg, Man. R3A 1R9, Canada. cbernst@cc.umanitoba.ca

SO Canadian Journal of Infectious Diseases, (1998) 9/5 (277-280).

Refs: 15

ISSN: 1180-2332 CODEN: CJDIES

CY Canada

DT Journal; Article

FS 004 Microbiology

005 General Pathology and Pathological Anatomy

LA English

SL English, French

AB Infection with Helicobacter ***pylori*** has been established as an important risk factor for the development of peptic ulcer disease, ***gastritis*** and ***gastric*** caner. The diagnosis of H ***pylori*** infection can be established by invasive or noninvasive techniques. Two noninvasive enzyme ***immunoassays*** (EIAs) for ***antibody*** detection - HeliSal and ***Pylori*** State - were compared with histology. Both assays detect immunoglobulin (Ig) G directed against purified H ***pylori*** antigen. The test populations consisted of 104 consecutive patients scheduled for upper gastrointestinal endoscopy. Of these patients, 97 (93%) had symptoms compatible with peptic ulcer disease. Saliva and serum were collected simultaneously at the time of endoscopy. Salivary EIA had a sensitivity of 66%, specificity of 67%, positive predictive value of 67% and negative predictive value of 66% compared with the serum EIA, where the results were 98%, 48%, 64% and 96%, respectively. Although the salivary EIA is an appealing noninvasive test, it was not a sensitive and specific assay. The serum EIA also lacked specificity, but was highly sensitive with a good negative predictive value. Although a negative serum EIA rules out H ***pylori*** infection, a positive must be interpreted in the clinical context and confirmed with a more specific measure.

L11 ANSWER 67 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 1998057224 EMBASE

TI Comparison of rapid office-based serology with formal laboratory-based

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ELISA testing for diagnosis of Helicobacter ***pylori***
    ***gastritis***
AU Kroser J.A.; Faigel D.O.; Furth E.E.; Metz D.C.
CS Dr. D.C. Metz, Gastroenterology Division, 3 Ravdin Building, Univ. of
   Pennsylvania Medical Center, 3400 Spruce Street, Philadelphia, PA 19104,
   United States
SO Digestive Diseases and Sciences, (1998) 43/1 (103-108).
   Refs: 37
   ISSN: 0163-2116 CODEN: DDSCDJ
CY United States
DT Journal; Article
FS 004 Microbiology
   029
         Clinical Biochemistry
   048 Gastroenterology
LA English
SL English
AB Accurate and cost-effective diagnosis of Helicobacter ***pylori***
    ***gastritis*** has taken on major importance. Several serologic tests
   for the diagnosis of H. ***pylori*** infection are commercially
   available. We compared the performance of the FlexSure HP rapid IgG
    ***antibody*** test with the conventional HM-CAP ELISA to evaluate
   whether qualitative office-based serology is reliable enough to replace
   formal laboratory-based testing. We assessed H. ***pylori*** status by
   concordance in 100 consecutive patients with antral biopsy, rapid urease,
   and 1 .mu.Ci [14C]urea breath tests. Both ***antibody*** tests had
   good sensitivity and specificity (> 86%). Concordance between the two
    ***antibody*** tests occurred in 87/93 patients (94%). Based on our
   data, the office-based FlexSure HP performed equally well as the
   laboratory-based formal ELISA and may be a better choice for initial
   serologic diagnosis in untreated patients.
L11 ANSWER 68 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.
AN 97327436 EMBASE
DN 1997327436
TI Relationship of helicobacter ***pylori*** infection to several
   malignant and non-malignant gastrointestinal diseases.
AU Erkisi M.; Colakoglu S.; Koksal F.; Tuncer I.; Burgut R.; Karakose H.;
   Doran F.; Zorludemir S.
CS M. Erkisi, Gazipasa Bulvari, Talay Apt. Kat: 1, Adana, Turkey
SO Journal of Experimental and Clinical Cancer Research, (1997) 16/3
   (289-294).
  Refs: 21
   ISSN: 0392-9078 CODEN: JECRDN
CY Italy
DT Journal; Article
FS 004 Microbiology
  016
         Cancer
         Public Health, Social Medicine and Epidemiology
  017
  048
         Gastroenterology
LA English
SL English
AB The importance of the Helicobacter ***Pylori*** infection was
  investigated as a risk factor for several gastrointestinal diseases. In
  this study 203 patients with ***gastric*** cancer, 61 with peptic
  ulcus, 60 with ***gastritis*** and 100 asymptomatic control subjects
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were investigated. Serum samples were examined for IgC ***antibodies*** to H. ***pylori*** by enzyme linked ***immunoassay*** - tissue samples were stained for H. ***pylori*** by Wartin-Stary technique and by Giemsa for routine histopathology. H. ***pylori*** seropositivity was 58.1% in ***gastric*** cancer, 54% in peptic ulcus, 63.3% in ***gastritis*** and 27% in asymptomatic control group. There was a 10.1% discordance between the serum and tumor determinants in the seropositive group and 11.3% of discordance in the seronegative group. The frequency of H. ***pylori*** seropositivity was lowest in cardia tumors (22.7%) and highest in antral tumors (65.5%, p = 0.00002). H. ***pylori*** seropositivity was 29% in diffuse type of histology, 35% in mixed type and 79% in the intestinal type (p = 0.00000). In the ***gastric*** cancer patients the frequent use of salty food (p = 0.00001, OR = 6.4), excessive salt, pickled food (p = 0.0000, OR = 24.92) and proteins (p = 0.003, OR =0.45) were more significant than asymptomatic volunteers. In ***gastric*** cancer patients the frequent use of salty and pickled food were relevantly associated with H. ***pylori*** infection (p = 0.001). It was concluded that H. ***pylori*** infection could play a role in the pathogenesis of non-malignant gastrointestinal diseases which may be the precursor of carcinoma. However, other contributing factors to carcinogenesis must be investigated.

L11 ANSWER 69 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 97325229 EMBASE

DN 1997325229

TI Immunoglobulin A ***antibodies*** to Helicobacter ***pylori***.

AU Jaskowski T.D.; Martins T.B.; Hill H.R.; Litwin C.M.

CS T.D. Jaskowski, ARUPICEP, 500 Chipeta Way, Salt Lake City, UT 84108, United States

SO Journal of Clinical Microbiology, (1997) 35/11 (2999-3000).

Refs: 16

ISSN: 0095-1137 CODEN: JCMIDW

CY United States

DT Journal; Article

FS 004 Microbiology

026 Immunology, Serology and Transplantation

LA English

SL English

AB Serological testing for immunoglobulin G (IgG) ***antibodies*** to Helicobacter ***pylori*** has proven useful in supporting the diagnosis of infection with this organism, but the clinical value of IgA ***antibodies*** in H. ***pylori*** -related ***gastritis*** remains controversial. The purpose of our study was to determine the frequency of IgA-positive IgG-negative patients with symptoms of gastrointestinal (GI) disorders, thus assessing the clinical utility of IgA testing for H. ***pylori*** -related ***gastritis*** . It was found previously that the frequency of infected individuals in this category (IgA positive and IgG negative) is about 2%, but a large number of IgG-negative patients with GI disorders suggestive of H. ***pylori*** infection have not been investigated until now.

L11 ANSWER 70 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 97258518 EMBASE

DN 1997258518

TI Evaluation of salivary ***antibodies*** to detect infection with

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Helicobacter ***pylori***
AU Loeb M.B.; Riddell R.H.; James C.; Hunt R.; Smaill F.M.
CS Dr. F.M. Smaill, University Medical Centre, 1200 Main Street West,
   Hamilton, Ont. L8N 3Z5, Canada. smaill@mcmaster.ca
SO Canadian Journal of Gastroenterology, (1997) 11/5 (437-440).
   Refs: 22
   ISSN: 0835-7900 CODEN: CJGAEJ
CY Canada
DT Journal; Article
FS 004 Microbiology
   005
         General Pathology and Pathological Anatomy
   026
         Immunology, Serology and Transplantation
   029
         Clinical Biochemistry
   048
         Gastroenterology
LA English
SL English; French
AB Helicobacter ***pylori*** infection is an important cause of peptic
   ulcer disease and chronic ***gastritis*** . Infection with this
   bacterium stimulates the production of immunoglobulin (Ig) G
    ***antibody*** . Salivary IgG ***antibody*** tests to detect H
    ***pylori*** infection offer a convenient and noninvasive method of
   diagnosis. To evaluate an IgG salivary ***antibody*** kit, saliva was
   collected from 157 out-patients with dyspepsia referred for endoscopy to a
   tertiary centre. A salivary IgG ELISA ***antibody*** assay was
   performed using the Helisal Helicobacter ***pylori*** (IgG) assay kit,
   and at least four ***gastric*** biopsies were obtained. H
    ***pylori*** infection was confirmed by demonstration of the organism on
   Warthin-Starry silver stain (sensitivity 85%, specificity 55%). The
   prevalence of infection with H ***pylori*** was 30%. When the analysis
   was redone, excluding those treated with eradication therapy, the results
   were similar (sensitivity 86%, specificity 58%). The positive predictive
   value of the assay was 45% and the negative predictive value was 90%.
   Despite the ease of sampling, the assay used has limited diagnostic
   utility, lacking the predictive value to indicate which patients referred
   with dyspeptic symptoms to a tertiary care setting are infected with H
    ***pylori*** .
L11 ANSWER 71 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.
AN 96371022 EMBASE
DN 1996371022
TI Evaluation of whole blood ***antibody*** kit to detect active
  Helicobacter ***pylori*** infection.
AU Borody T.J.; Andrews P.; Shortis N.P.
CS Centre for Digestive Diseases, 144 Great North Road, Five Dock, NSW 2046,
   Australia
SO American Journal of Gastroenterology, (1996) 91/12 (2509-2512).
  ISSN: 0002-9270 CODEN: AJGAAR
CY United States
DT Journal; Article
FS 004
         Microbiology
  005
         General Pathology and Pathological Anatomy
  026
         Immunology, Serology and Transplantation
  037
         Drug Literature Index
         Gastroenterology
  048
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LA English

SL English

AB Objectives: To evaluate the sensitivity and specificity of a whole blood ***antibody*** test (Helisal(TM) Rapid Blood test) for the detection of Helicobacter ***pylori*** using endoscopic diagnostic criteria of histology and urease tests as the 'gold standard.' Methods: A prospective trial of Helisal(TM) Rapid Blood (HRB) test was carried out in patients undergoing investigations for dyspepsia that included endoscopic biopsy for rapid urease test, microbiological culture, and histology. Blood samples were obtained at the time of endoscopy and were tested for the presence of ***antibody*** to H. ***pylori*** using the HRB test. In a separate patient group, results of ***antibody*** tests in whole venous and capillary blood were compared (n = 25). Results: The rapid blood test was carried out immediately after the endoscopic examination with a result available in under 10 min in all cases. In 203 patients examined, the HRB test detected 70 of 203 to be H. ***pylori*** positive as compared with 71 of 203 using urease/histology. Against combined urease/histology tests, the HRB test achieved 82% sensitivity and 91% specificity. Five patients were judged to be 'false negative' on endoscopic tests for H. ***pylori*** (extensive intestinal metaplasia n = 3; recent use of antimicrobials) yet the HRB test diagnosed the presence of infection, which could be shown to resolve on treatment. The HRB achieved 89% sensitivity and 91% specificity upon correctly including these five patients in the calculations. In all 25 patients tested, venous and capillary blood results concurred giving HRB test positivity in each case. Conclusions: Whether using whole venous or capillary blood, the HRB test is a quick, convenient, and accurate test for the diagnosis of active H. ***pylori*** infection in patients previously not treated. In a subgroup of patients with low level infection due to recent antimicrobials or intestinal metaplasia negative to all endoscopic tests, the blood test can still correctly diagnose H. ***pylori*** infection. Because blood samples require no centrifugation before testing, the greatest usefulness of this test will be that of a primary office diagnostic device.

L11 ANSWER 72 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 96150640 EMBASE

DN 1996150640

TI Comparison of rapid serological tests (FlexSure HP and QuickVue) with conventional ELISA for detection of Helicobacter ***pylori*** infection.

AU Graham D.Y.; Evans Jr. D.J.; Peacock J.; Baker J.T.; Schrier W.H.

CS Veterans Affairs Medical Center, 2002 Holcombe Blvd., Houston, TX 77030, United States

SO American Journal of Gastroenterology, (1996) 91/5 (942-948). ISSN: 0002-9270 CODEN: AJGAAR

CY United States

DT Journal; Article

FS 004 Microbiology

006 Internal Medicine

048 Gastroenterology

LA English

SL English

AB Background: There is a need for accurate and rapid tests for Helicobacter

pylori infection especially since the recent National Institutes
of Health Consensus Development Conference on H. ***pylori*** in
peptic ulcer disease charged the medical community with treating H.

pylori infection in all patients with H. ***pylori*** and ulcer disease. Methods: We prospectively compared a simple, rapid serological test (FlexSure HP, SmithKline Diagnostics) for the detection of serum IgG ***antibodies*** against H. ***pylori*** with another rapid test (QuickVue, Quidel) and two enzyme ***immunoassays*** (HM-CAP, Enteric Products, and PyloriStat, BioWhittaker). Serum samples from 551 individuals including both symptomatic patients (196) and asymptomatic volunteers (355) were tested for the presence of IgG ***antibodies*** against H. ***pylori*** . The presence or absence of active H. ***pylori*** infections was determined using the [14C]-urea breath test. Results: All of the serological tests performed well. FlexSure HP had calculated sensitivity, specificity, and accuracy of 94.4, 87.6, and 91.1%, respectively, relative to the urea breath test. In 49 of the 551 samples, the urea breath test and FlexSure HP did not agree. Those samples were tested with HM-CAP ***immunoassay*** to confirm presence or absence of IgG ***antibodies*** against H. ***pylori*** . After the resolution of the discordant results, the sensitivity, specificity, and accuracy of FlexSure HP were 96.0, 95.1, and 95.6%, respectively, and were comparable to HM-CAP and PyloriStat. FlexSure HP was compared with histology or culture in 75 cases, and the accuracy was 100%. FlexSure HP and QuickVue were compared using 200 serum samples. FlexSure HP was more specific (88.7 vs 79.4%) and accurate (91 vs 84%) than QuickVue (p < 0.05 for both), relative to the urea breath test with discordant samples unresolved. FlexSure HP was also simpler to use, easier to interpret, and faster than QuickVue. FlexSure HP required no sample dilution, one reagent, four steps, and 5 min to complete. Conclusion: FlexSure HP is an excellent option for in-office tests for the physician who desires immediate results or for small laboratories that do not have the volume of H. ***pylori*** testing to justify ELISA test formats.

L11 ANSWER 73 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 96053180 EMBASE

DN 1996053180

TI Helicobacter ***pylori*** and recurrent abdominal pain in children.

AU Hardikar W.; Feekery C.; Smith A.; Oberklaid F.; Grimwood K.

CS Department of Pediatrics, Yale University School of Medicine, 333 Cedar Street, New Haven, CT 06510, United States

SO Journal of Pediatric Gastroenterology and Nutrition, (1996) 22/2 (148-152).

ISSN: 0277-2116 CODEN: JPGND6

CY United States

DT Journal; Article

FS 007 Pediatrics and Pediatric Surgery

048 Gastroenterology

LA English

SL English

AB Recurrent abdominal pain is one of the most common presentations to pediatricians; yet an organic etiology can be found in only 10% of cases. Because infection with Helicobacter ***pylori*** in adults and children results in ***gastritis***, a causative role for the organism has been postulated. To investigate this theory, we conducted a prospective case-control study in children with recurrent abdominal pain using serum H. ***pylori*** IgG ***antibodies*** measured by an enzyme immunoabsorbent assay. Age, sex, ethnicity, and socioeconomic status were adjusted in the statistical model. Five subjects (5.1%) and 14

controls (14.3%) had raised serum IgG ***antibodies*** to H.

pylori (adjusted OR, 0.21; 95% confidence interval, 0.05, 0.85).

The negative association between H. ***pylori*** and recurrent abdominal pain indicates that this organism is unlikely to have an important etiologic role in this disorder.

L11 ANSWER 74 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 95318303 EMBASE

DN 1995318303

TI A practical single sample dry latex agglutination test for Helicobacter

pylori ***antibody*** detection.

AU Midolo P.D.; Lambert J.R.; Russell E.G.; Lin S.K.

CS Department of Microbiology, Monash Medical Centre, 246 Clayton Road, Clayton 3168, Vic., Australia

SO Journal of Clinical Pathology, (1995) 48/10 (969-971). ISSN: 0021-9746 CODEN: JCPAAK

CY United Kingdom

DT Journal; Article

FS 004 Microbiology

005 General Pathology and Pathological Anatomy

048 Gastroenterology

LA English

SL English

AB Assessment of a single serum sample for Helicobacter ***pylori*** ***antibodies*** is frequently requested in routine diagnostic laboratories. Current enzyme linked immunosorbent assay (ELISA) kits are not ideal for testing small numbers of serum samples and some have low sensitivities, specificities or large grey zones.-A panel of 90 serum samples from patients who had presented for routine upper endoscopy was used to compare three kits for the detection of H ***pylori*** ***antibodies*** : (1) Pyloriset Dry, total ***antibody*** latex agglutination, Orion Diagnostica, Espoo, Finland; (2) Pyloriset enzyme ***immunoassay*** (EIA), IgG ELISA, Orion; and (3) Hel-p, IgG ELISA, Amrad, Kew, Victoria, Australia. Diagnosis of H ***pylori*** positivity was made if culture results and either rapid urease test or histopathology were positive. The sensitivity, specificity, positive, and negative predictive value for each test was as follows: Orion: latex 93.3%, 95.6%, 95.5%, 93.3%, respectively; Orion: EIA-G 84.4%, 97.8%, 97.4%, 84.4%, respectively; and Amrad: EIA-G 100%, 88.9%, 90%, 100%, respectively. The latex test performed better than the EIAs with respect to sensitivity and specificity.

L11 ANSWER 75 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 93326585 EMBASE

DN 1993326585

TI Prevalence of immunoglobulin G ***antibodies*** to Helicobacter ***pylori*** in Chilean individuals [7].

AU Figueroa G.; Troncoso M.; Portell D.P.; Toledo M.S.; Acuna R.; Arellano L.

CS Microbiology Unit, Inst Nutrition and Food Technology, University of Chile, Casilla 138-11, Santiago, Chile

SO European Journal of Clinical Microbiology and Infectious Diseases, (1993) 12/10 (795-797).

ISSN: 0934-9723 CODEN: EJCDEU

CY Germany

DT Journal; Letter

FS 004 Microbiology 017 Public Health, Social Medicine and Epidemiology LA English L11 ANSWER 76 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. AN 93258309 EMBASE DN 1993258309 TI Use of serum specific immunoglobulin ***antibodies*** to determine helicobacter ***pylori*** associated ***gastritis*** AU Fischbach W.; Wosnik K.; Kirchner T.; Mossner J. CS Medizinische Poliklinik, University of Wurzburg, Klinikstr. 8,D-97070 Wurzburg, Germany SO Zeitschrift für Gastroenterologie, (1993) 31/7-8 (429-431). ISSN: 0044-2771 CODEN: ZGASAX CY Germany DT Journal; Article FS 004 Microbiology 020 Gerontology and Geriatrics Immunology, Serology and Transplantation 026 048 Gastroenterology LA English SL English; German AB A prospective study in 169 consecutive patients referred for upper gastrointestinal endoscopy was initiated to investigate the diagnostic performance of serum Helicobacter ***pylori*** (HP) specific immunoglobulin (Ig) G ***antibodies*** Using an enzyme linked immunosorbent assay (ELISA) an excellent correlation between serologic evidence of HP and the demonstration of this organism by histology and urease test in 79 H P-positive patients was found. Serum IgG also correlated with the histological degree and the activity of ***gastritis*** . Our results demonstrate that serum IG G ***antibodies***, as determined by ELISA, are highly useful for diagnosis of HP-associated ***gastritis*** .

L11 ANSWER 77 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 91315671 EMBASE

DN 1991315671

TI High prevalence of Helicobacter ***pylori*** infection and histologic ***gastritis*** in asymptomatic hispanics.

AU Dehesa M.; Dooley C.P.; Cohen H.; Fitzgibbons P.L.; Perez-Perez G.I.; Blaser M.J.

CS Depts. Medicine and Pathology, Univ. of Southern California, School of Medicine, Los Angeles, CA 90033, United States

SO Journal of Clinical Microbiology, (1991) 29/6 (1128-1131). ISSN: 0095-1137 CODEN: JCMIDW

CY United States

DT Journal; Article

FS 004 Microbiology

005 General Pathology and Pathological Anatomy

026 Immunology, Serology and Transplantation

048 Gastroenterology

LA English

SL English

AB In this study, we estimated the prevalence of Helicobacter ***pylori***
infection and histologic ***gastritis*** in 58 asymptomatic Hispanic

adult volunteers (mean age, 41 years; 59% male) by endoscopic biopsy of the upper gastrointestinal tract. Forty-six subjects (79%) were found to harbor H. ***pylori*** in ***gastric*** biopsies, and all had histologic ***gastritis***. Four other subjects were found to have ***gastritis*** in the absence of H. ***pylori***. Similar prevalences of H. ***pylori*** and ***gastritis*** were noted in all age groups and also in American-born and immigrant Hispanics. Biopsy data and serologic studies of H. ***pylori*** ***antibodies*** correlated well. We conclude that H. ***pylori*** infection is an almost universal finding in the ***gastric*** mucosa of asymptomatic adult Hispanics, regardless of age. The clinical significance of these findings is unknown, but we speculate that H. ***pylori*** and its associated ***gastritis*** could have a role in the high incidence of ****gastric*** carcinoma in Hispanic populations.

L11 ANSWER 78 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 91312270 EMBASE

DN 1991312270

- TI A solid-phase enzyme-linked immunospot (ELISPOT) assay for detection of Helicobacter ***pylori*** ***antibody*** -producing cells in ***gastric*** mucosa.
- AU Sugiyama T.; Furuyama S.; Awakawa T.; Imai K.; Yabana T.; Yachi A.; Yokota Oguma K.K.
- CS Dept. of Internal Medicine, Sapporo Medical College, S-l, W-16, Chuo-ku, Sapporo 060, Japan
- SO Gastroenterologia Japonica, (1991) 26/5 (684). ISSN: 0435-1339 CODEN: GAJABC

CY Japan

DT Journal; Article

FS 004 Microbiology

026 Immunology, Serology and Transplantation

048 Gastroenterology

LA English

- L11 ANSWER 79 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.
- AN 90034384 EMBASE
- DN 1990034384
- TI Value of serology (ELISA and immunoblotting) for the diagnosis of Campylobacter ***pylori*** infection.
- AU Pena A.S.; Endtz Ph. H.; Offerhaus G.J.A.; Hoogenboom-Verdegaal A.; Van Duijn W.; De Vargas N.; Den Hartog G.; Kreuning J.; Van der Reyden J.; Mouton R.P.; Lamers C.B.H.W.
- CS Department of Gastroenterology, Leiden University Hospital, P.O. Box 9600,NL-2300 RC Leiden, Netherlands
- SO Digestion, (1989) 44/3 (131-141).

ISSN: 0012-2823 CODEN: DIGEBW

CY Switzerland

DT Journal; Article

FS 029 Clinical Biochemistry

048 Gastroenterology

- LA · English
- SL English
- AB Fifty-two unselected patients referred to for upper gastrointestinal endoscopy were evaluated in several ways to determine the presence of Campylobacter ***pylori*** . ***Antibodies*** against this

microorganism were measured to assess the value of serology for the diagnosis of C. ***pylori*** infection. Five antral biopsy specimens were taken in each patient for culture and bacteriological determinations, histology [morphology and Warthin-Starry (WS) staining] and the urease test (2, 3 and 24 h). Serum ***antibodies*** against a sonicate of 6 strains of microorganisms were assayed by enzyme-linked ***immunoassay*** (ELISA) and an immunoblotting technique. In 14 of the 52 patients the histology of the antrum was normal, 18 patients had chronic active ***gastritis*** and 20 had chronic ***gastritis*** without polymorphonuclear infiltration. In the group with normal histology, only 1 patient was positive for C. pilory with all methods, and 1 other subject was positive for IgG and 2 for IgA only with ELISA. In the group with chronic active ***gastritis***, 14 were positive with all methods, 1 was negative by WS only and another was negative for IgA according to ELISA, WS and ***antibodies*** . Among the patients with chronic ***gastritis***, 7 were positive and 7 negative with all tests; in the other 6 patients the results obtained with the various tests were divergent. Four serological tests were studied and validated against culture, WS and urease test which were considered to be the reference methods. The serological tests showed high sensitivity and specificity for the detection of C. ***pylori*** -associated active chronic ***gastritis*** of the antrum, and can therefore serve as noninvasive methods to identify individuals with this condition.

L11 ANSWER 80 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 89163233 EMBASE

DN 1989163233

TI Serum IgG and IgA ***antibody*** responses to campylobacter ***pylori*** in a group of healthy asymptomatic volunteers.

AU Westblom T.U.; Barthel J.S.; Kosunen T.U.; Everett E.D.

CS Department of Medicine, Section of Infectious Diseases, Marshall University School of Medicine, Huntington, WV 25755-9410, United States

SO Scandinavian Journal of Infectious Diseases, (1989) 21/3 (311-314). ISSN: 0036-5548 CODEN: SJIDB7

CY Sweden

DT Journal

FS 004 Microbiology

026 Immunology, Serology and Transplantation

048 Gastroenterology

LA English

SL English

AB Sera from 17 healthy asymptomatic volunteers were tested for presence of IgG and IgA ***antibodies*** against Campylobacter ***pylori*** and correlated with endoscopic biopsy findings. Three volunteers infected with C. ***pylori*** had the highest IgG ***antibody*** titers of the group. None of 14 C. ***pylori*** free subjects had significant IgG ***antibody*** levels. IgA ***antibody*** titers were negative in all subjects regardless of state of infection, in contrast to control sera from symptomatic C. ***pylori*** infected patients who manifested high IgA ***antibody*** levels.

L11 ANSWER 81 OF 184 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 89071885 EMBASE

DN 1989071885

TI Age-dependent increase of Campylobacter ***pylori***

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***antibodies*** in blood donors.
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- AU Kosunen T.U.; Hook J.; Rautelin H.I.; Myllyla G.
- CS Department of Bacteriology and Immunology, University of Helsinki, 00290 Helsinki, Finland
- SO Scandinavian Journal of Gastroenterology, (1989) 24/1 (110-114). ISSN: 0036-5521 CODEN: SJGRA4
- CY Norway
- DT Journal
- FS 048 Gastroenterology
- LA English
- SL English
- AB ***Antibodies*** against Campylobacter ***pylori*** were determined in 500 blood donors aged 18 to 65 years. Acid extract from a C. ***pylori*** strain was used as antigen in enzyme ***immunoassay*** The proportion of donors with high ***antibody*** titers increased with age. For IgG ***antibodies*** it was 10% in the age group from 18 to 25 years but 60% in the group from 56 to 65 years; the increase for IgA and IgM ***antibodies*** was from 5 to 42% and from 7 to 21%, respectively. The geometric mean titers of those with high values showed no clear changes with age, which would imply chronic antigenic stimulus.
- L11 ANSWER 82 OF 184 LIFESCI COPYRIGHT 2001 CSA
- AN 97:17620 LIFESCI
- TI Campylobacter ***pylori*** antigens and uses thereof for detection of Campylobacter ***pylori*** infection
- CS ENTERIC RESEARCH LABORATORIES, INC.
- SO (1995) . US Patent 5459041; US Cl. 435/7.21 435/7.3 435/7.92 435/7.93 435/7.94 435/7.95 435/961 435/974 435/975 436/518 436/527 436/528 436/529 436/531 436/533 436/547 436/804 530/350 530/.
- DT Patent
- FS A; W3
- LA English
- AB Antigenic compositions are disclosed for use in diagnostic kits and the like for detecting the presence of ***antibodies*** specific for Campylobacter ***pylori***, bacteria often associated with the occurrence of Type B ***gastritis*** and peptic ulcer disease. Samples of bodily fluids, for instance, may be contacted with immobilized antigen which is then washed and tested for the occurrence of significant levels of antigen/ ***antibody*** complex. Levels exceeding a predetermined positive threshold are indicative of ***antibodies*** to Campylobacter ***pylori*** in the sample tested. Kits employing the antigenic compositions of the invention preferably include means for detecting the antigen/ ***antibody*** complex such as materials and reagents for conducting an enzyme-linked immunosorbent assay, Western blot technique, liposome-based assay or other known detection tests.
- L11 ANSWER 83 OF 184 LIFESCI COPYRIGHT 2001 CSA
- AN 89:107753 LIFESCI
- TI Time-resolved fluoroimmunoassay for Campylobacter ***pylori***
 antibodies.
- AU Aceti, A.; Pennica, A.; Leri, O.; Caferro, M.; Grilli, A.; Celestino, D.; Casale, V.; Citarda, F.; Grassi, A.; Sciarretta, F.
- CS Inst. Trop. and Infect. Dis., La Sapienza Univ., 00161 Rome, Italy
- SO LANCET., (1989) vol. 2, no. 8661, p. 505.
- DT Journal

FS J; F

LA English

AB Dr. Loffeld and colleagues suggest that an ELISA test for Campylobacter

pylori ***antibodies*** might replace endoscopy in the
diagnosis of ***gastritis*** associated with this bacterium. However,
with a cut-off of optical density (OD) greater than 2 multiplied by 1 the
ELISA had a specificity of 100% and a sensitivity of 85 multiplied by 4%;
at a lower cut-off (OD above 1) the sensitivity was 100% but the
specificity fell to 72 multiplied by 7%. We have evaluated a time-resolved
fluoroimmunoassay (TR-FIA), to detect C. ***pylori***

antibodies This test is based on the labelling of

antibodies with europium (Eu) and conversion of the specifically
bound non-fluorescent label to highly fluorescent chelate solution,
followed by measurement with a time-resolved fluorimeter. TR-FIA was
compared with ELISA.

L11 ANSWER 84 OF 184 MEDLINE

AN 2000451530 MEDLINE

DN 20460284

- TI Accuracy of an enzyme ***immunoassay*** for the detection of Helicobacter ***pylori*** in stool specimens in the diagnosis of infection and posttreatment check-up.
- AU Forne M; Dominguez J; Fernandez-Banares F; Lite J; Esteve M; Gali N; Espinos J C; Quintana S; Viver J M
- CS Department of Gastroenterology, Hospital Universitari Mutua de Terrassa, Barcelona, Spain.
- SO AMERICAN JOURNAL OF GASTROENTEROLOGY, (2000 Sep) 95 (9) 2200-5. Journal code: 3HE. ISSN: 0002-9270.
- CY United States
- DT Journal; Article; (JOURNAL ARTICLE)
- LA English
- FS Priority Journals
- EM 200012
- EW 20001203
- AB OBJECTIVE: The aim of this study was to assess the reliability of a newly developed enzyme ***immunoassay*** for Helicobacter ***pylori*** -specific antigen detection in stools (HpSA) compared to other standardized diagnostic techniques such as histology (H), rapid urease test (RUT) and 13C-urea breath test (UBT) to diagnose H. ***pylori*** infection and to evaluate its usefulness in determining H. ***pylori*** status after treatment. METHODS: One hundred eighty-eight patients referred to our department for upper gastrointestinal endoscopy were included. H. ***pylori*** infection was confirmed in all patients by HpSA test in stools, RUT, UBT, and H. Patients were defined as positive for H. ***pylori*** if RUT and UBT or H were positive. A total of 142 symptomatic patients received eradication treatment and were reassessed 6 wk after therapy; for 70 of these patients, stool samples were also collected at 24 h and 6 months after finishing eradication treatment. In the posttreatment follow-up, UBT was used as gold standard. RESULTS: The sensitivity of HpSA test for the diagnosis of H. ***pylori*** infection using a cut-off value of 0.130 was 89.5% and its specificity 77.8%. This specificity was lower than that obtained with UBT, H, and RUT. In the early follow-up the sensitivity of HpSA test was null. At 6 weeks and at 6 months post-treatment its sensitivity was 70.4% and 50% and its specificity was 81.6% and 79.3%, respectively. CONCLUSIONS: The HpSA stool

test, using a cut-off value of 0.130, may be useful for the primary diagnosis of H. ***pylori*** infection, with sensitivity similar to that obtained with other standard tests, but with less specificity. HpSA test is not useful for early monitoring of treatment efficacy. At 6 wk and at 6 months posttreatment, HpSA test lacks accuracy as compared to UBT for evaluating the outcome of the eradication treatment.

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L11 ANSWER 85 OF 184 MEDLINE
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AN 2000386919 MEDLINE

DN 20319644

- TI Study of diagnostic modalities and pathology of Helicobacter
 pylori infection in children.
- AU Bansal D; Patwari A K; Logani K B; Malhotra V L; Anand V K
- CS Department of Pediatrics, Lady Hardinge Medical College, New Delhi.
- SO INDIAN JOURNAL OF PATHOLOGY AND MICROBIOLOGY, (1999 Jul) 42 (3) 311-5. Journal code: GKK. ISSN: 0377-4929.
- CY India
- DT Journal; Article; (JOURNAL ARTICLE)
- LA English
- EM 200010
- EW 20001002
- AB To evaluate various diagnostic tests for Helicobacter ***pylori*** (Hp) in children, and to study the spectrum of endoscopic and histological changes in the stomach and duodenum of children with gastroduodenal disorders, associated with Hp infection Children below 12 years of age with various gastroduodenal disorders requiring upper gastrointestinal endoscopy were studied. Endoscopic biopsy specimens were collected from duodenum and antrum. Apart from histopathological examination of biopsy material, rapid urease test (RUT) of the antral biopsy specimen and blood examination to estimate specific IgG ***antibodies*** to Hp by Indirect Solid Phase Enzyme ***Immunoassay*** was performed. Forty seven children were included. Nine (19.1%) of them were positive both by serology and RUT. Seven (14.9%) were positive by histology. A significant correlation of Hp was noticed with chronic antral ***gastritis*** (p = 0.002) and chronic duodenitis (p = 0.006). Age equal to or more than 10 years was found to be significant risk factor for acquiring Hp infection. Prevalence of Hp in children with gastroduodenal complaints was found to be 19%. Both RUT and serology were found to be reliable diagnostic tests for Hp as compared with histology. Antral ***gastritis*** and chronic duodenitis had a significant correlation with Hp colonization.
- L11 ANSWER 86 OF 184 MEDLINE
- AN 2000282994 MEDLINE
- DN 20282994
- TI Plaunotol suppresses interleukin-8 secretion induced by Helicobacter

 pylori : therapeutic effect of plaunotol on H. ***pylori***
 infection.
- AU Takagi A; Koga Y; Aiba Y; Kabir A M; Watanabe S; Ohta-Tada U; Osaki T; Kamiya S; Miwa T
- CS Department of Internal Medicine, Tokai University School of Medicine, Isehara, Kanagawa, Japan.. takagia@is-icc.u-tokai.ac.jp
- SO JOURNAL OF GASTROENTEROLOGY AND HEPATOLOGY, (2000 Apr) 15 (4) 374-80. Journal code: A6J. ISSN: 0815-9319.
- CY Australia
- DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 200009

EW 20000905

AB BACKGROUND: It has been suggested that ***gastric*** mucosal injury induced by Helicobacter ***pylori*** infection is mediated by interleukin-8 (IL-8). METHODS: We studied the effect of plaunotol, a drug extracted from the Plau-noi tree of Thailand, and reported it to be effective in the treatment of ulcers, of IL-8 secretion induced by H. ***pylori*** and of the inhibitory adhesion activity of the bacterium to ***gastric*** epithelial cells. Moreover, the therapeutic effect of plaunotol on H. ***pylori*** infection was assessed by using the gnotobiotic murine model. RESULTS: Plaunotol inhibited the growth of H. ***pylori*** (1.5 x 10(4) c.f.u./mL) at high doses (24-48 microg/mL), but not at low doses (3-6 microg/mL). Interleukin-8 secretion induced by H. ***pylori*** was inhibited by coculture with plaunotol in a dose-dependent manner. The adhesion of H. ***pylori*** to MKN45 cells was also suppressed by coculture with plaunotol in a dose-dependent manner. An in vivo study showed that plaunotol improved histological ***gastritis*** and decreased the H. ***pylori*** ***antibody*** titre. CONCLUSIONS: These findings suggest that plaunotol has a therapeutic effect on ***gastritis*** induced by H. ***pylori***.

L11 ANSWER 87 OF 184 MEDLINE

AN 2000131489 MEDLINE

DN 20131489

TI Evaluation of an enzyme ***immunoassay*** for detecting Helicobacter ***pylori*** antigens in human stool samples.

AU Agha-Amiri K; Mainz D; Peitz U; Kahl S; Leodolter A; Malfertheiner P

CS Department of Gastroenterology, Hepatology and Infectious Diseases, Otto-von-Guericke-University, Magdeburg, Germany.

SO ZEITSCHRIFT FUR GASTROENTEROLOGIE, (1999 Dec) 37 (12) 1145-9. Journal code: XU1. ISSN: 0044-2771.

CY GERMANY: Germany, Federal Republic of

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 200005

EW 20000501

AB BACKGROUND AND AIM: So far, the detection of Helicobacter ***pylori***

(Hp) infection by stool analysis appeared to be almost impossible. With
the Premier Platinum HpSA EIA a new enzyme ***immunoassay*** was
developed for diagnosis of Hp infection, using polyclonal

antibodies against Hp antigens in human stool. We evaluated this new test in its diagnostic accuracy in comparison to established reference methods. METHODS: From 54 consecutive patients (29 male, 25 female, age: 19 to 85 years) undergoing routine upper gastrointestinal endoscopy antral and corpus biopsies were taken for histology and Helicobacter urease test (HUT). Endoscopy, 13C-urea breath test (13C-UBT), serology, and stool probes sampling were performed within two days. Stool samples were aliquoted after reception and stored frozen (-20 degrees C) until tested. The Premier Platinum HpSA test (Meridian, Connecticut, Ohio, USA) was performed according to the manufactures protocol. Patients were considered to be infected with Hp if two of the four reference tests were positive. RESULTS: 28 of the 54 patients were Hp-infected. Only one of these was

found to be false-negative by the HpSA EIA. Two false-positive results were obtained in the noninfected group (sensitivity 96.4%, specificity 92.3%). CONCLUSION: In this group of patients investigated, the novel HpSA Enzyme ***Immunoassay*** (EIA) proved to be highly accurate for diagnosis of Hp infection. Collection and testing of stool are noninvasive and easy to perform, therefore this test will become an important tool for diagnosing Hp infection in clinical practice.

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L11 ANSWER 88 OF 184 MEDLINE
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AN 2000033670 MEDLINE

DN 20033670

TI Evaluation of three commercial serological tests with different methodologies to assess Helicobacter ***pylori*** infection.

AU van Der Ende A; van Der Hulst R W; Roorda P; Tytgat G N; Dankert J

CS Department of Medical Microbiology, Amsterdam, The Netherlands.

SO JOURNAL OF CLINICAL MICROBIOLOGY, (1999 Dec) 37 (12) 4150-2. Journal code: HSH. ISSN: 0095-1137.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 200003

EW 20000303

AB The sera of 142 Helicobacter ***pylori*** -positive and 32 H.

pylori -negative patients were assessed by a desktop test

(Quick Vue), an enzyme-linked immunosorbent assay (ELISA) (HM-CAP), and a solid-phase, two-step chemiluminescent enzyme ***immunoassay***

(Immulite). These tests yielded sensitivities of 97, 97, and 91% and specificities of 97, 94, and 100%, respectively. In conclusion, the desktop test and the ELISA are more sensitive than the chemiluminescent enzyme ***immunoassay*** (P < 0.05). The chemiluminescent enzyme

immunoassay has the advantage that it is fully automated.

L11 ANSWER 89 OF 184 MEDLINE

AN 1999405995 MEDLINE

DN 99405995

TI Helicobacter ***pylori*** ***antibody*** profile in household members of children with H. ***pylori*** infection.

AU Elitsur Y; Adkins L; Saeed D; Neace C

CS Department of Pediatrics, Marshall University, School of Medicine, Huntington, WV 25701-0195, USA.

SO JOURNAL OF CLINICAL GASTROENTEROLOGY, (1999 Sep) 29 (2) 178-82. Journal code: IBG. ISSN: 0192-0790.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 199912

EW 19991204

AB Intrafamilial spread is implicated as a major route for acquisition of Helicoobacter ***pylori*** infection. Investigating H. ***pylori*** cytotoxin-associated protein (CagA) and vacuolating toxin (VacA) ***antibodies*** within family members enabled the authors to evaluate this possibility further. Serum samples were collected prospectively from household members after their index children were diagnosed with active H.

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***pylori*** infection. Serum samples were evaluated for anti-H.
 ***pylori*** immunoglobulin G ***antibody*** using the enzyme
 ***immunoassay*** (IEA) method and for H. ***pylori*** CagA and VacA
 ***antibodies*** with the commercially available immunoprobing Western
blot kit. Ten different families participated in the study, including 10
pediatric patients and 31 household members. All patients and 28 household
members (90%) were seropositive for H. ***pylori*** ***antibody***
by IEA and Western blot tests. Overall, 17 subjects (41.4%) were CagA
positive, 14 (34.1%) were VacA positive, 11 (26.8%) were positive for both
 ***antibodies*** , and 22 (53.6%) were negative for both ***antibodies*** . A significant association in bacterial
 ***antibody*** profile was found between the patient index members and
all household members (Cohen's kappa and Mentel-Haenszel methods). In four
families, more than 66% of the household members harbored the same
 ***antibody*** profile, and in two families a completely different
profile was observed. Moreover, a similar H. ***pylori***
 ***antibody*** profile between the index patient and the mother was
found in six families, and between the index patient and the father in two
families. The data strongly suggest an intrafamiliar transmission for H.
 ***pylori*** infection.
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L11 ANSWER 90 OF 184 MEDLINE

AN 1999388378 MEDLINE

DN 99388378

TI New immunological assays for the diagnosis of Helicobacter ***pylori***
infection

AU Vaira D; Holton J; Menegatti M; Ricci C; Landi F; Ali' A; Gatta L; Acciardi C; Farinelli S; Crosatti M; Berardi S; Miglioli M

CS Department of Internal Medicine, University of Bologna, Bologna, Italy.

SO GUT, (1999 Jul) 45 Suppl 1 I23-7. Ref: 51 Journal code: FVT. ISSN: 0017-5749.

CY ENGLAND: United Kingdom

DT Journal; Article; (JOURNAL ARTICLE)
General Review; (REVIEW)
(REVIEW, TUTORIAL)

LA English

FS Abridged Index Medicus Journals; Priority Journals

EM 199911

EW 19991105

AB There are several types of immunological tests available for the diagnosis and management of Helicobacter ***pylori*** infection. Most commercially available serological kits use the enzyme linked immunosorbent assay (ELISA) test format. Originally the kits used crude antigen preparations although many of the newer kits use a more purified antigen preparation, with often increased specificity but lower sensitivity. Near patient test kits are based either on latex agglutination or immunochromatography. Generally they have low sensitivities compared with laboratory tests. Western blotting, ELISA, and recombinant immunoblot assays (RIBA) have also been developed into commercially available kits and can be used to indicate the presence of specific virulence markers. An antigen detection kit has been developed for the detection of Helicobacter ***pylori*** in faeces.

Immunological reagents have also been combined with other diagnostic modalities to develop immunohistochemical stains and DNA

immunoassays . Helicobacter ***pylori*** is now recognised as

the cause of ***gastritis*** and most cases of peptic ulcer disease (PUD); its long term carriage increases the risk of ***gastric*** adenocarcinoma sixfold and it is designated as a class I carcinogen. H ***pylori*** has also been implicated as a cause of ***gastric*** mucosa associated lymphoid tissue lymphomas. Its relation to non-ulcer dyspepsia remains controversial. Additionally, long term carriage of the organism may be associated with short stature in young girls and, in the general population, as a possible risk factor for the development of vasospastic disorders and possibly skin immunopathology such as urticaria. With the recognition of H ***pylori*** as an important human pathogen, it has become one of the growing number of organisms to have its complete genome sequence mapped. Serology is an important method of determining colonisation status and can be used for diagnosis, as a screening procedure, or to follow the efficacy of eradication regimens. Most serological assays are in the ELISA format although some are based on the latex agglutination reaction. These latter are used principally as near patient assays. Most assays detect IgG in serum although some detect serum IgA. More recently developed assays detect IgA in saliva and the production of affinity purified ***antibodies*** has led to the development of an antigen detection assay for faecal specimens. Serological reagents have also been used in immunocytochemistry and to speed up the detection of amplified products of the polymerase chain reaction (PCR)-DNA ***immunoassays***

L11 ANSWER 91 OF 184 MEDLINE

AN 1999359814 MEDLINE

DN 99359814

TI Long-term follow-up study of serum immunoglobulin G and immunoglobulin A ***antibodies*** after Helicobacter ***pylori*** eradication.

AU Kato S; Furuyama N; Ozawa K; Ohnuma K; Iinuma K

CS Department of Pediatrics, Tohoku University School of Medicine, Japan.. skato@ped.med.tohoku.ac.jp

SO PEDIATRICS, (1999 Aug) 104 (2) e22. Journal code: CZE. ISSN: 1098-4275.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 199910

EW 19991003

AB OBJECTIVE: There have been few studies concerning serum titers of anti-Helicobacter ***pylori*** immunoglobulin G (IgG) ***antibody*** >12 months after eradication of the original infection. Moreover, clinical usefulness of immunoglobulin A (IgA) ***antibody*** levels remains to be established. The purpose of this study was to investigate long-term responses of serum IgG-specific and IgA-specific ***antibodies*** to H ***pylori*** in children after eradication therapy. STUDY DESIGN: A total of 34 children, 2 to 17 years of age (mean: 11.7 years) with H ***pylori*** -associated gastroduodenal disease received eradication therapy (proton pump inhibitor-based dual or triple regimens). Diagnoses included nodular ***gastritis*** (n = 8), ***gastric*** ulcer (n = 7), and duodenal ulcer (n = 19). Upper gastrointestinal endoscopy and biopsy were performed before the therapy and at 1 to 2 months' posttreatment. H ***pylori*** infection and eradication were defined by biopsy-based tests; eradication was successful in 28 patients and

unsuccessful in 6. Pretreatment IgG was positive in 30 patients (88. 2%), and the IgA was positive in 31 (91.2%), who were entered into this study (duration </=24 months). Serum samples were obtained before treatment and at 1, 3, 6, 12, 18, and 24 months' posttreatment. IgG and IgA ***antibodies*** were measured using commercial enzyme ***immunoassay*** kits (HM-CAP and PP-CAP; Enteric Products, Inc., New York, NY). RESULTS: Compared with pretreatment values, IgG and IgA ***antibodies*** significantly and steadily decreased at 1 through 24 months' posttreatment in successfully treated patients. A decrease in titer of the IgA class was significantly greater than that of the IgG class at 1 to 12 months' follow-up. There was no significant decrease in titer of either ***antibody*** in all but 2 patients with eradication failure. A >/=30% decrease in titer of the IgA ***antibody*** at 6 months indicated eradication with sensitivity of 90.5% and specificity of 100%. For the IgG ***antibody***, a 30% decrease at 12 months showed equal sensitivity and specificity. Seroreversion rates of IgG and IgA ***antibodies*** were 53% and 48% at 12 months and were 86% and 81% at 24 months, respectively. The mean periods from the completion of eradication therapy to seroreversion of IgG and IgA ***antibodies*** were 11.2 +/- 7.0 and 11.6 +/- 7.8 months, respectively (not significantly different). A higher pretreatment titer of IgG ***antibody*** was related to a longer period of seroreversion (r = 0.44). In one patient, (13)C-urea breath test-confirmed reinfection was accompanied by reappearance of significant titers of the IgG and IgA ***antibodies*** . CONCLUSIONS: A serology test is useful for evaluating eradication in children. Approximately half of patients with successful eradication remained to be IgG-seropositive and IgA-seropositive at 12 months' posttreatment. When a decrease titer in ***antibody*** is used for assessing eradication, an endpoint of >/=6 months is required. The IgA ***antibody*** may be a more convenient indicator of H ***pylori*** status than is the IgG ***antibody*** .

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L11 ANSWER 92 OF 184 MEDLINE
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AN 1999311132 MEDLINE

DN 99311132

TI Prevalence of CagA, VacA ***antibodies*** in symptomatic and asymptomatic children with Helicobacter ***pylori*** infection.

AU Elitsur Y; Neace C; Werthammer M C; Triest W E

CS Department of Pediatrics, Marshall University School of Medicine, Huntington, West Virginia 25701-0195, USA.

SO HELICOBACTER, (1999 Jun) 4 (2) 100-5. Journal code: CY4. ISSN: 1083-4389.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 199910

EW 19991001

AB BACKGROUND: Limited data are available on the prevalence of CagA and VacA
Helicobacter ***pylori*** ***antibodies*** in children. The aim of
this study was to investigate the ***antibody*** prevalence to the H.

pylori virulence factors CagA and VacA in symptomatic and
asymptomatic children with H. ***pylori*** infection and to correlate
these ***antibodies*** with the severity of ***gastric***
inflammation or density of H. ***pylori*** organisms in the

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***gastric*** mucosa. MATERIALS AND METHODS: Twenty-three symptomatic
   children and 132 asymptomatic children with positive H. ***pylori***
   serology participated in this study. Anti-H. ***pylori*** IgG
    ***antibody*** and CagA or VacA H. ***pylori***
                                                        ***antibodies***
   were measured by enzyme ***immunoassay*** (HM-CAP; sensitivity and
   specificity > 90%) and Western immunoblot (Helicoblot 2.0) methods,
   respectively. ***Gastric*** inflammation and H. ***pylori***
   density were graded histologically using the revised Sydney criteria.
   RESULTS: The prevalence of CagA and VacA ***antibodies*** were 69% and
   35% in symptomatic children and 54% and 52% in asymptomatic children.
   respectively. Multiple regression analysis showed a correlation between
   CagA ***antibody*** and the severity of ***gastritis*** but no
   correlation with other histological features, including the number of
   neutrophils or lymphoid follicles. Neither ***antibody*** correlated
  with the degree of bacterial density in the ***gastric*** mucosa.
   CONCLUSION: CagA and VacA H. ***pylori*** ***antibodies*** are
   common in the pediatric population. The combined CagA/VacA
    ***antibodies*** correlated weakly with the degree of mucosal
  inflammation.
L11 ANSWER 93 OF 184 MEDLINE
AN 1999129624 MEDLINE
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DN 99129624

TI Helicobacter ***pylori*** infection in children with celiac disease: prevalence and clinicopathologic features.

AU Luzza F; Mancuso M; Imeneo M; Mesuraca L; Contaldo A; Giancotti L; La Vecchia A M; Docimo C; Pensabene L; Strisciuglio P; Pallone F; Guandalini

CS Dipartimento di Medicina Sperimentale e Clinica, Universit'a di Catanzaro Magna Graecia, Italy.

SO JOURNAL OF PEDIATRIC GASTROENTEROLOGY AND NUTRITION, (1999 Feb) 28 (2) 143-6.

Journal code: JL6. ISSN: 0277-2116.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 199906

EW 19990601

AB BACKGROUND: Celiac disease is frequently associated with chronic ***gastritis*** . Helicobacter ***pylori*** is the main etiologic agent of chronic ***gastritis*** . The aim of this study was to assess the prevalence of H. ***pylori***, the related symptoms, and the endoscopic and histologic ***gastric*** features in children with celiac disease. METHODS: Eight-one (24 boys, 57 girls; age range: 1.4-17.7 years, median 6.8) children with celiac disease were studied. All children had a blood sample taken. In a subgroup of 30 children who underwent endoscopy, three ***gastric*** biopsy specimens were taken for histology (hematoxylin and eosin, Giemsa, immunohistochemistry) and urease quick test. Symptom complaints were recorded. Age- and sex-matched (one case, one control) children without celiac disease were used for comparison. Serum H. ***pylori*** IgG were measured by means of a locally validated commercial enzyme-linked ***immunoassay*** . RESULTS: Overall, 15 of 81 (18.5%) children with celiac disease and 14 of 81 (17.3%) control children were positive for H. ***pylori*** . The

percentage of H. ***pylori*** positivity was similar in children with untreated and treated celiac disease. Recurrent abdominal pain was the only symptom that helped to distinguish between H. ***pylori***
-positive and H. ***pylori*** -negative children. However, symptoms disappeared in patients with celiac disease after gluten withdrawal, irrespective of H. ***pylori*** status. All endoscopic (erythema, nodularity) and histologic (superficial-, interstitial-, lymphocytic***gastritis***, activity, lymphoid follicles) findings did not differ between celiac and nonceliac H. ***pylori*** -positive children.
CONCLUSIONS: Prevalence and clinical expressivity of H. ***pylori*** infection is not increased in children with celiac disease. The clinicopathologic pattern of the infection is not specifically influenced in this condition.

L11 ANSWER 94 OF 184 MEDLINE

AN 1998443826 MEDLINE

DN 98443826

TI Helicobacter ***pylori*** infection in recurrent abdominal pain.

AU Bansal D; Patwari A K; Malhotra V L; Malhotra V; Anand V K

CS Department of Pediatrics and Microbiology, Lady Hardinge Medical College, New Delhi.

SO INDIAN PEDIATRICS, (1998 Apr) 35 (4) 329-35. Journal code: GM2. ISSN: 0019-6061.

CY India

DT (CLINICAL TRIAL)

(CONTROLLED CLINICAL TRIAL)

Journal; Article; (JOURNAL ARTICLE)

LA English

EM 199901

EW 19990104

AB OBJECTIVE: To study the relationship between Helicobacter ***pylori*** (Hp) infection and recurrent abdominal pain (RAP) and to evaluate various modalities to diagnose Hp infection. DESIGN: Prospective case control study. SETTING: Teaching hospital. METHODS: Children between 3-12 years of age with RAP in whom upper gastrointestinal endoscopic examination was indicated were studied. Endoscopic biopsy specimen were collected from duodenum, antrum and esophagus. Apart from histopathological examination of biopsy material, rapid urease test (RUT) of the antral biopsy specimen and blood examination to estimate specific IgG ***antibodies*** to Hp by Indirect Solid Phase Enzyme ***Immunoassay*** was performed. The results of Hp IgG ***antibodies*** was compared with age matched controls. RESULTS: Thirty one children with RAP were subjected to endoscopic examination and their anti Hp IgG ***antibodies*** status compared with 26 controls. Hp colonization was detected in 7 children (23%) with RAP; by RUT in 23% and antral biopsy in 16% of cases. Anti Hp IgG ***antibodies*** were also positive in almost equal proportion (19%) of controls (p = 0.757). Endoscopic examination revealed esophagitis in 16% of cases and none had evidence of ***gastric*** or duodenal erosion, ulcer or cobblestone appearance of antrum. A significant correlation of Hp was noticed with chronic antral ***gastritis*** (p = 0.002), chronic duodenitis (p = 0.02) and age > 10 years (p = 0.02). No significant correlation was noticed between Hp colonization and various socioeconomic risk factors. CONCLUSION: Hp does not seem to be commonly associated with RAP in our patient population as Hp colonization was detected in only 23% of cases which was not significantly higher than the

seroprevalence of anti Hp IgG ****antibodies*** in the controls. However, a small sample size of our study limits drawing any firm conclusions. Antral ***gastritis*** and chronic duodenitis had a significant correlation with Hp colonization. RUT was found to be a reliable diagnostic test to detect Hp.

L11 ANSWER 95 OF 184 MEDLINE

AN 1998403644 MEDLINE

DN 98403644

TI Helicobacter ***pylori*** infection: an added stressor on iron status of women in the community.

AU Peach H G; Bath N E; Farish S J

CS University of Melbourne, Ballarat Health Services Base Hospital, VIC.. a.temperley@gpph.unimelb.edu.au

SO MEDICAL JOURNAL OF AUSTRALIA, (1998 Aug 17) 169 (4) 188-90. Journal code: M26. ISSN: 0025-729X.

CY Australia

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals; Cancer Journals

EM 199811

EW 19981103

AB OBJECTIVE: To explore a possible association between Helicobacter

pylori infection and iron status. DESIGN: Cross-sectional study.

SETTING: Ballarat (a major regional city in Victoria), population 78000,
October November 1997. PARTICIPANTS: 160 women and 152 men, a subsample of
participants in a cardiovascular disease risk factor prevalence survey for
whom frozen plasma was available. MAIN OUTCOME MEASURES: H. ***pylori***

IgG ***antibody*** status by enzyme ***immunoassay***; iron
intake; plasma iron, transferrin and ferritin concentrations. RESULTS: 28%
of women and 33% of men were infected with H. ***pylori***. The mean
(SEM) plasma ferritin concentration of infected women (59.3 [7.6]
microg/L) was significantly lower than for non-infected women (88.8 [7.9]
microg/L; P=0.002), after adjusting for age. Mean daily dietary iron
intakes were similar in infected and non-infected women. CONCLUSIONS: H.

pylori infection appears to be an additional stressor on women's
iron status, but the mechanism remains to be determined.

L11 ANSWER 96 OF 184 MEDLINE

AN 1998081473 MEDLINE

DN 98081473

TI Comparison between a rapid office-based and ELISA serologic test in screening for Helicobacter ***pylori*** in children.

AU Elitsur Y; Neace C; Triest W E

CS Department of Pediatrics, Marshall University School of Medicine, Huntington, WV 25701-0195, USA.. yelitsur@musom.marshall.edu

SO HELICOBACTER, (1997 Dec) 2 (4) 180-4. Journal code: CY4. ISSN: 1083-4389.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 199803

EW 19980304

AB BACKGROUND. The rapid diagnostic serological test for detection of

Helicobacter ***pylori*** (H. ***pylori***) infection in children has a significant advantage over the standard enzyme ***immunoassay*** (EIA) method, for its simplicity and rapid availability of results in a physician's office setting. We compared the immunochromatographic test with a standard enzyme ***immunoassay*** test in the pediatric population. MATERIALS AND METHODS. A retrospective analysis of 1147 serum samples from asymptomatic children and prospective analysis of 62 serum samples from symptomatic children undergoing diagnostic upper endoscopy were evaluated for the detection of H. ***pylori*** ***antibodv*** by two commercially available serology tests. Each serum sample was tested by a rapid test (FlexSure HP, SmithKline Diagnostics, Inc.) and compared to the standard EIA method (HM-CAP, Enteric Products, Inc.). RESULTS. The rapid test, FlexSure HP, was comparable to the rapid EIA test in screening for H. ***pylori*** infection in symptomatic and asymptomatic children with sensitivity and specificity of 83-90% and 90-100%, respectively. Both methods had a comparable sensitivity and specificity for the detection of H. ***pylori*** -associated ***gastritis*** (60-70% and 94%. respectively). CONCLUSION. The rapid test is comparable to the standard EIA test and may be used by physicians in symptomatic children. The use of FlexSure HP as a screening tool for the prevalence of H. ***pylori*** infection in asymptomatic children may be limited by its low positive predictive value compared to the EIA method.

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L11 ANSWER 97 OF 184 MEDLINE
AN 97268485 MEDLINE
DN 97268485
TI [The pathogenic role of Helicobacter ***pylori***].
  O patogennoi roli Helicobacter ***pylori*** .
AU Ivashkin V T; Polozhentsev S D; Sultanov V K; Blinov V D; Solomakhin S V;
  Kretsu A P; Kalinin V K; Spesivtsev V N
SO TERAPEVTICHESKII ARKHIV, (1993) 65 (2) 11-3.
  Journal code: VLU. ISSN: 0040-3660.
CY RUSSIA: Russian Federation
DT Journal; Article; (JOURNAL ARTICLE)
LA Russian
FS Priority Journals
EM 199707
EW 19970701
AB The urease test and bacterioscopy of impression smears were used to detect
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Helicobacter ***pylori*** (HP) in biopsies from pyloric

gastric mucosa of 77 (89.5%) of 86 chronic ***gastritis***

patients, of 27 (4%) from 32 duodenal ulcer patients, of 84 (84.0%) from

100 healthy male subjects aged 18-20. There was focal hyperemia in pyloric
part of the stomach in 59 patients, leukocytic infiltration of mucous
membrane was found histologically in 78 ones. Close correlation between
complaints, endoscopic and histological shifts, HP incidence rate was not
registered. Positive results in determination of HP ***antibodies***
by enzyme ***immunoassay*** (EIA) were obtained in 32 (45.7%) of
healthy subjects. EIA findings and histological evidence on HP presence
failed to coincide in 42 (92.9%) duodenal ulcer patients.

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L11 ANSWER 98 OF 184 MEDLINE
AN 97259999 MEDLINE
DN 97259999
TI Prevalence of Helicobacter ***pylori*** ***antibodies*** in
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children in Bloemfontein, South Africa. AU Pelser H H; Househam K C; Joubert G; van der Linde G; Kraaij P; Meinardi M; McLeod A; Anthony M CS Department of Paediatrics and Child Health, University of the Orange Free State, Bloemfontein, South Africa. SO JOURNAL OF PEDIATRIC GASTROENTEROLOGY AND NUTRITION, (1997 Feb) 24 (2) 135-9. Journal code: JL6. ISSN: 0277-2116. CY United States DT Journal; Article; (JOURNAL ARTICLE) LA English FS Priority Journals EM 199709 EW 19970903 AB BACKGROUND: An association of H. ***pylori*** infection with chronic ***gastritis***, peptic ulceration and ***gastric*** cancer is known. METHODS: Prevalence of IgG ***antibodies*** to Helicobacter ***pylori*** in children in the Bloemfontein, South Africa area was studied. Children attending the general pediatric outpatient department at Pelonomi Hospital in Bloemfontein were grouped according to age. A minimum of 100 children was investigated in each age group. Baseline demographic and socioeconomic data were collected. RESULTS: The study showed a high prevalence of H. ***pylori*** ***antibodies*** Prevalence increased with age: 13.5% in children 3 months-2 years, 48.5% at 2-5 years, 67.3% at 5-10 years and 84.2% at 10-15 years. Investigation of the socioeconomic data in relation to the prevalence of H. ***pylori*** was inconclusive. CONCLUSIONS: This high prevalence needs further study. L11 ANSWER 99 OF 184 MEDLINE AN 93213900 MEDLINE DN 93213900 TI [Evaluation of the cut-off point in the serological diagnosis of Helicobacter ***pylori*** infection in children using an enzyme ***immunoassay*** technique (letter)]. Valoracion del punto de corte (cut off) en el diagnostico serologico de la infeccion por Helicobacter ***pylori*** en ninos mediante una tecnica de enzimoinmunoanalisis. AU Sanz J C; Martin E; Alarcon T; Martinez M J; Garcia-Novo M D; Lopez-Brea M SO ENFERMEDADES INFECCIOSAS Y MICROBIOLOGIA CLINICA, (1993 Jan) 11 (1) 55. Journal code: A10. ISSN: 0213-005X. CY Spain DT Letter LA Spanish EM 199307 L11 ANSWER 100 OF 184 MEDLINE AN 91041503 MEDLINE DN 91041503 TI [Detection of ***antibodies*** to Helicobacter ***pylori*** with the immunoenzyme test and indirect immunofluorescence]. Nachweis von Antikorpern gegen Helicobacter ***pylori*** mit Enzymimmuntest und indirekter Immunfluoreszenz. AU Abb J; Striegel K; Fruhmorgen P CS Mikrobiologisches Institut, Krankenanstalten Ludwigsburg..

SO LEBER, MAGEN, DARM, (1990 Sep) 20 (5) 224-30.

Journal code: L3P. ISSN: 0300-8622.

CY GERMANY: Germany, Federal Republic of DT Journal; Article; (JOURNAL ARTICLE)

LA German

FS Priority Journals

EM 199102

AB Sera from 56 adult patients were screened for the presence of IgG ***antibodies*** against Helicobacter ***pylori*** by enzyme ***immunoassay*** and indirect immunofluorescence. In addition, the detection of Helicobacter ***pylori*** in antral biopsy specimens was attempted by culture and histological methods. Colonisation of the antrum mucosa with Helicobacter ***pylori*** was observed in 39 of the 56 patients. IgG ***antibodies*** against Helicobacter ***pylori*** were detected by enzyme ***immunoassay*** in 34 of 39 infected patients. Thus, the enzyme ***immunoassay*** showed a sensitivity of 87.2 percent and a specificity of 82.4 percent. IgG ***antibodies*** against Helicobacter ***pylori*** were further detected by indirect immunofluorescence in 28 of 39 infected patients. Thus, indirect immunofluorescence showed a sensitivity of 66.7 percent and a specificity of 88.2 percent. Our results demonstrate that the enzyme ***immunoassay*** for IgG ***antibodies*** and other invasive or noninvasive methods for the detection of infection with Helicobacter ***pylori*** appear to be of equal sensitivity and specificity.

L11 ANSWER 101 OF 184 MEDLINE

AN 88257555 MEDLINE

DN 88257555

TI Immunoblot analysis of immune response to Campylobacter ***pylori*** and its clinical associations.

'AU von Wulffen H; Grote H J; Gatermann S; Loning T; Berger B; Buhl C

CS Institut für Medizinische Mikrobiologie und Immunologie, Universitätskrankenhaus Eppendorf, Hamburg, Federal Republic of Germany...

SO JOURNAL OF CLINICAL PATHOLOGY, (1988 Jun) 41 (6) 653-9. Journal code: HT3. ISSN: 0021-9746.

CY ENGLAND: United Kingdom

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Abridged Index Medicus Journals; Priority Journals; Cancer Journals

EM 198810

AB Systemic immune response to Campylobacter ***pylori*** was detected by the immunoblot technique in serum samples from 200 patients, 129 blood donors, and 96 children. The results of the IgG immunoblot test showed excellent correlation with the detection of C ***pylori*** by culture and also with histopathological examination of the antrum, as well as with peptic ulcer disease. An IgA response also occurred and gave results comparable with those of the IgG immunoblot test, although on a quantitatively lower scale. The IgM immunoblots were of no help in the serodiagnosis of C ***pylori*** infection. The protein bands that seemed to be the most specific for C ***pylori*** and which were consistently observed in patients positive for C ***pylori*** were a 110 kilodalton and a 63 kilodalton band on the IgG immunoblot and an 89 kilodalton band on the IgA immunoblot. A 94 kilodalton and a 28 kilodalton band were also included in the evaluation. While immunoblot analysis may be used effectively for the serodiagnosis of C ***pylori*** infection and can distinguish between patients with normal antrum mucosa and those

with ***gastritis***, the test does not help to distinguish between those patients with antrum ***gastritis*** who subsequently develop peptic ulcers and those who do not.

L11 ANSWER 102 OF 184 MEDLINE

AN 88149892 MEDLINE

DN 88149892

TI Campylobacter ***pylori*** in Swedish patients referred for gastroscopy.

AU Gnarpe H; Unge P; Blomqvist C; Makitalo S

CS Department of Clinical Bacteriology, Gavle Central Hospital, Sweden...

SO APMIS, (1988 Feb) 96 (2) 128-32. Journal code: AMS. ISSN: 0903-4641.

CY Denmark

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals; Cancer Journals

EM 198806

AB Campylobacter ***pylori*** was isolated more often from patient with peptic ulcers and in an age-related manner in a material of 395 consecutive patients referred for gastroscopy. Direct microscopy was done with Gram and Acridine Orange stain and found too insensitive for practical use. All patients were investigated serologically with an enzyme ***immunoassay*** which showed excellent correlation with positive cultures for C. ***pylori***. The findings were discussed, and the enzyme ***immunoassay*** , with a negative predictive value of 0.99, was found to be a valuable tool for primary screening of patients suspected of being carriers of C. ***pylori***.

L11 ANSWER 103 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 2000:79951 SCISEARCH

GA The Genuine Article (R) Number: 276BQ

TI Seroepidemiology of Helicobacter ***pylori*** infection in a Jamaican community

AU Lindo J F; LynSue A E; Palmer C J; Lee M G; Vogel P; Robinson R D (Reprint)

CS UNIV W INDIES, DEPT LIFE SCI, KINGSTON 7, JAMAICA (Reprint); UNIV W INDIES, DEPT LIFE SCI, KINGSTON 7, JAMAICA; UNIV W INDIES, DEPT MICROBIOL, KINGSTON 7, JAMAICA; UNIV MIAMI, SCH MED, CTR DIS PREVENT, MIAMI, FL; UNIV W INDIES, DEPT MED, KINGSTON 7, JAMAICA

CYA JAMAICA; USA

SO TROPICAL MEDICINE & INTERNATIONAL HEALTH, (DEC 1999) Vol. 4, No. 12, pp. 862-866.

Publisher: BLACKWELL SCIENCE LTD, P O BOX 88, OSNEY MEAD, OXFORD OX2 0NE, OXON, ENGLAND.

ISSN: 1360-2276.

DT Article; Journal

FS CLIN

LA English

REC Reference Count: 40

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB We researched epidemiologic associations between environmental and demographic factors and prevalence of Helicobacter ***pylori*** infection in a suburban Jamaican community. Using a clustered sampling technique, 22 domestic yards enclosing 60 separate households were

randomly selected from a local community. All household members (n = 346)were invited to participate following informed consent, the overall compliance rate was 58.9%. A commercial enzyme ***immunoassay*** (HMaCAP) was used to detect IgG ***antibodies*** raised against H. ***pylori*** . Environmental and demographic information was obtained by questionnaire. The seroprevalence of H. ***pylori*** was 69.9% (n = 202). Analysis of the independent variables revealed three major components: Component 1 described, collectively, good personal hygiene and sanitation, indoor water supply and absence of straying animals in the peridomestic area; Component 2 included older age, good personal hygiene and large yard size; Component 3 the presence of domestic animals (cats and dogs) and, again, large vard size. These three complexes explained 42.2% of the variability in the data set. Logistic regression showed that Components 2 and 3 were independently associated with H. ***pylori*** seropositivity, indicating that a combination of demographic, environmental and zoonotic factors is involved in the spread of H. ***pylori*** infections at the tropical community level.

L11 ANSWER 104 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 1999:613049 SCISEARCH

GA The Genuine Article (R) Number: 222EP

TI Long-term follow-up study of serum immunoglobulin G and immunoglobulin A
antibodies after Helicobacter ***pylori*** eradication

AU Kato S (Reprint); Furuyama N; Ozawa K; Ohnuma K; Iinuma K

CS TOHOKU UNIV, SCH MED, DEPT PEDIAT, AOBA KU, 1-1 SEIRYO MACHI, SENDAI, MIYAGI 9808574, JAPAN (Reprint); SENDAI CITY HOSP, DEPT PEDIAT, SENDAI, MIYAGI, JAPAN

CYA JAPAN

SO PEDIATRICS, (AUG 1999) Vol. 104, No. 2, Part 1, pp. E221-E225.
Publisher: AMER ACAD PEDIATRICS, 141 NORTH-WEST POINT BLVD, ELK GROVE VILLAGE, IL 60007-1098.
ISSN: 0031-4005.

15514. 0051-4005.

DT Article; Journal

FS LIFE; CLIN

LA English

REC Reference Count: 25

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB Objective. There have been few studies concerning serum titers of anti-Helicobacter ***pylori*** immunoglobulin G (IgG) ***antibody*** >12 months after eradication of the original infection. Moreover, clinical usefulness of immunoglobulin A (IgA) ***antibody*** levels remains to be established. The purpose of this study was to investigate long-term responses of serum IgG-specific and IgA-specific ***antibodies*** to H ***pylori*** in children after eradication therapy.

Study Design. A total of 34 children, 2 to 17 years of age (mean: 11.7 years) with H ***pylori*** -associated gastroduodenal disease received eradication therapy (proton pump inhibitor-based dual or triple regimens). Diagnoses included nodular ***gastritis*** (n = 8), ***gastric*** ulcer (n = 7), and duodenal ulcer (n = 19). Upper gastrointestinal endoscopy and biopsy were performed before the therapy and at 1 to 2 months' posttreatment. H ***pylori*** infection and eradication were defined by biopsy-based tests; eradication was successful in 28 patients and unsuccessful in 6. Pretreatment IgG was positive in 30 patients (88.2%), and the IgA was positive in 31 (91.2%), who were entered into this study (duration less than or equal to 24 months). Serum samples were

obtained before treatment and at 1, 3, 6, 12, 18, and 24 months' posttreatment. IgG and IgA ***antibodies*** were measured using commercial enzyme ***immunoassay*** kits (HM-CAP and PP-CAP; Enteric Products, Inc, New York, NY).

Results. Compared with pretreatment values, IgG and IgA ***antibodies*** significantly and steadily decreased at 1 through 24 months' posttreatment in successfully treated patients. A decrease in titer of the IgA class was significantly greater than that of the IgG class at 1 to 12 months' follow-up. There was no significant decrease in titer of either ***antibody*** in all but 2 patients with eradication failure. A greater than or equal to 30% decrease in titer of the IgA ***antibody*** at 6 months indicated eradication with sensitivity of 90.5% and specificity of 100%. For the IgG ***antibody***, a 30% decrease at 12 months showed equal sensitivity and specificity. Seroreversion rates of IgG and IgA ***antibodies*** were 53%; and 48% at 12 months and were 86% and 81% at 24 months, respectively. The mean periods from the completion of eradication therapy to seroreversion of IgG and IgA ***antibodies*** were 11.2 +/- 7.0 and 11.6 +/- 7.8 months, respectively (not significantly different). A higher pretreatment titer of Ige ***antibody*** was related to a longer period of seroreversion (r = 0.44). In one patient, C-13-urea breath test-confirmed reinfection was accompanied by reappearance of significant titers of the IgG and IgA ***antibodies*** .

Conclusions. A serology test is useful for evaluating eradication in children. Approximately half of patients with successful eradication remained to be IgG-seropositive and IgA-seropositive at 12 months' posttreatment When a decrease titer in ***antibody*** is used for assessing eradication, an endpoint of greater than or equal to 6 months is required. The IgA ***antibody*** may be a more convenient indicator of H ***pylori*** status than is the IgG ***antibody***

L11 ANSWER 105 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 1999:531148 SCISEARCH

GA The Genuine Article (R) Number: 212NW

TI Helicobacter ***pylori*** in the Canadian arctic: Seroprevalence and detection in community water samples

AU McKeown I; Orr P; Macdonald S; Kabani A; Brown R; Coghlan G; Dawood M; Embil J; Sargent M; Smart G; Bernstein C N (Reprint)

CS GB443 HLTH SCI CTR, GASTROENTEROL SECT, 820 SHERBROOK ST, WINNIPEG, MB R3A 1R9, CANADA (Reprint), UNIV MANITOBA, DEPT MED, WINNIPEG, MB, CANADA, UNIV MANITOBA, DEPT COMMUNITY HLTH SCI, WINNIPEG, MB R3T 2N2, CANADA; UNIV MANITOBA, DEPT MED MICROBIOL, WINNIPEG, MB, CANADA; NWT, KEEWATIN REG HLTH BOARD, WINNIPEG, MB, CANADA; CADHAM PROV LAB, RH RES LAB, WINNIPEG, MB, CANADA

CYA CANADA_

SO AMERICAN JOURNAL OF GASTROENTEROLOGY, (JUL-1999)-Vol.-94, No.-7, pp. 1823-1829.

Publisher: ELSEVIER SCIENCE INC, 655 AVENUE OF THE AMERICAS, NEW YORK, NY 10010.

ISSN: 0002-9270.

DT Article; Journal

FS CLIN

LA English

REC Reference Count: 25

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB OBJECTIVE: Many North American arctic communities are characterized by risk markers associated with Helicobacter ***pylori*** (H. ***pylori***) infection, including overcrowded housing and inadequate water supply and sanitation systems. Our aim was to determine the seroprevalence of H. ***pylori*** infection in two traditional Inuit communities in the central Canadian arctic and to test for the presence of H. ***pylori***, by polymerase chain reaction (PCR), in local water supplies.

METHODS: Samples of venous whole blood from adults and capillary blood

from children were collected and analyzed by enzyme ***immunoassay*** and Helisal Rapid Test, respectively, for IgG ***antibody*** to H.

pylori ***Antibodies*** to CagA were detected by enzyme

immunoassay, and ABO and Lewis antigens were also determined.

Demographic and clinical information were collected by questionnaire.

Water samples from each community were tested for H. ***pylori*** by

PCR.

RESULTS: One hundred-thirty (50.8%) of 256 subjects from the two communities were positive for H. ***pylori*** IgG ***antibodies***. Seropositive subjects were more likely to be male, compared with seronegative individuals (p = 0.01). ***Antibody*** status did not differ with respect to age, community, alcohol or cigarette use, number of persons per household, gastrointestinal complaints or previous investigations, medications, or presence of blood group O, Lewis a-b+. CagA ***antibodies*** were detected in 78 (61.9%) of 126 H. ***pylori*** -seropositive subjects tested; however, 41 (35.3%) of 116 H. ***pylori*** -seronegative subjects were also CagA positive. Water samples taken from the water delivery truck in Chesterfield Inlet and two lakes near Repulse Bay were positive for H. ***pylori*** in the study group was higher than rates in southern Canadian populations, but lower

CONCLUSION: The seroprevalence of H. ***pylori*** in the study group was higher than rates in southern Canadian populations, but lower than the seroprevalence previously documented in a Canadian subarctic Indian (First Nations) community. The detection of H. ***pylori*** in local water supplies may indicate a natural reservoir for the organism or possible contamination from human sewage. (Am J Gastroenterol 1999;94:1823-1829. (C) 1999 by Am. Coll. of Gastroenterology).

L11 ANSWER 106 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 1998:719184 SCISEARCH

GA The Genuine Article (R) Number: 119LD

TI Evaluation of commercially available Helicobacter ***pylori*** serology kits: a review

AU Laheij R J F (Reprint); Straatman H; Jansen J B M J; Verbeek A L M

CS MIES 152, POB 9101, NL-6500 HB NIJMEGEN, NETHERLANDS (Reprint); UNIV NIJMEGEN HOSP, DEPT GASTROENTEROL, NL-6500 HB NIJMEGEN, NETHERLANDS; UNIV NIJMEGEN, DEPT EPIDEMIOL, NIJMEGEN, NETHERLANDS
CYA NETHERLANDS

SO JOURNAL OF CLINICAL MICROBIOLOGY, (OCT 1998) Vol. 36, No. 10, pp. 2803-2809.

Publisher: AMER SOC MICROBIOLOGY, 1325 MASSACHUSETTS AVENUE, NW, WASHINGTON, DC 20005-4171.

ISSN: 0095-1137.

DT General Review; Journal

FS LIFE, CLIN

LA English

REC Reference Count: 90

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AN 1998:699502 SCISEARCH
GA The Genuine Article (R) Number: 117NF
 TI Analysis of immunoglobulin a ***antibodies*** to Helicobacter
    ***pylori*** in serum and ***gastric*** juice in relation to mucosal
   inflammation
AU Hayashi S (Reprint); Sugiyama T; Yokota K; Isogai H; Isogai E; Oguma K;
   Asaka M; Fujii N; Hirai Y
CS JICHI MED SCH, DEPT MICROBIOL, 3311-1 YAKUSHIJI, MINAMI KAWACHI, TOCHIGI
   3290498, JAPAN (Reprint); HOKKAIDO UNIV, SCH MED, DEPT INTERNAL MED 3,
   SAPPORO, HOKKAIDO 060863, JAPAN; OKAYAMA UNIV, SCH MED, DEPT BACTERIOL,
   OKAYAMA 7008558, JAPAN; SAPPORO MED UNIV, SCH MED, DEPT MICROBIOL,
   SAPPORO, HOKKAIDO 060855, JAPAN; SAPPORO MED UNIV, SCH MED, ANIM
   EXPERIMENTAT CTR, SAPPORO, HOKKAIDO 060855, JAPAN
CYA JAPAN
SO CLINICAL AND DIAGNOSTIC LABORATORY IMMUNOLOGY, (SEP 1998) Vol. 5, No. 5,
   pp. 617-621.
   Publisher: AMER SOC MICROBIOLOGY, 1325 MASSACHUSETTS AVENUE, NW.
   WASHINGTON, DC 20005-4171.
   ISSN: 1071-412X.
DT Article: Journal
FS LIFE
LA English
REC Reference Count: 33
   *ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS*
      Helicobacter ***pylori*** is a major etiologic agent in
   gastroduodenal disorders. In this study, immunoglobulin A (IgA)
    ***antibodies*** to H. ***pylori*** antigens were evaluated in serum
   and ***gastric*** juice specimens obtained from patients with
    ***gastritis*** or peptic ulcers by utilizing ***antibody*** capture
   enzyme-linked immunosorbent assays (ACELISAs). Urease alpha subunit (UA),
   urease beta subunit (UB), the 66-kDa heat shock protein (HSP), and the
   25-kDa protein (25K) were used as antigens for the ACELISAs. The
    ***antibody*** titers of the ACELISAs reflect the ratio of H.
    ***pylori*** -specific IgA to total IgA. The ratio is stable, although
   the ***antibody*** concentration fluctuates in ***gastric***
  juice. By using ACELISAs it was possible to evaluate quantitatively not
  only serum IgA ***antibodies*** but also ***gastric*** juice
   secretory IgA (S-IgA) ***antibodies*** . In both serum IgA and
   ***gastric*** juice S-IgA ACELISAs, the titers of ***antibody*** to
  HSP and 25K were remarkably correlated with the histologic grade of
    ***gastritis***, whereas those to UA and UB were not strongly correlated
  with histologic grade. Thus, it is useful for estimating the histologic
   grade of ***gastritis*** to quantify serum IgA and ***gastric***
  juice S-IgA ***antibodies*** to HSP and 25K.
L11 ANSWER 108 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)
AN 97:774500 SCISEARCH
GA The Genuine Article (R) Number: YB027
TI Helicobacter ***pylori*** Lewis expression is related to the host
  Lewis phenotype
AU Wirth H P (Reprint); Yang M Q; Peek R M; Tham K T; Blaser M J
CS UNIV ZURICH, SCH MED, DIV GASTROENTEROL, RAMISTR 100, CH-8091 ZURICH,
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SWITZERLAND (Reprint); VANDERBILT UNIV, SCH MED, DIV INFECT DIS.

L11 ANSWER 107 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

NASHVILLE, TN 37212; VANDERBILT UNIV, SCH MED, DIV GASTROENTEROL, NASHVILLE, TN 37212; DEPT VET AFFAIRS MED CTR, NASHVILLE, TN 37212 CYA SWITZERLAND: USA

SO GASTROENTEROLOGY, (OCT 1997) Vol. 113, No. 4, pp. 1091-1098.
Publisher: W B SAUNDERS CO, INDEPENDENCE SQUARE WEST CURTIS CENTER, STE 300, PHILADELPHIA, PA 19106-3399.
ISSN: 0016-5085.

DT Article; Journal

FS LIFE; CLIN

LA English

REC Reference Count: 35

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

Background & Aims: Lewis antigens occur in human ***gastric*** epithelium and in Helicobacter ***pylori*** lipopolysaccharide; their expression is polymorphic in both. Autoimmune mechanisms induced by bacterial Lewis expression have been proposed to cause ***gastritis*** . The aim of this study was to examine the relationship between bacterial and host ***gastric*** Lewis expression, as determined by the erythrocyte Lewis(a/b) phenotype, and between ***gastric*** histopathology and bacterial Lewis expression. Methods: H. ***pylori*** Lewis expression was determined by enzyme ***immunoassays***, erythrocyte Lewis phenotype was assessed by agglutination tests, and ***gastric*** histopathology was scored blindly. Results: The host Lewis phenotype was (a+b-) in 15, (a-b+) in 34, and (a-b-) in 17 patients, therefore expressing Lewis x, y, or neither as their major ***gastric*** epithelial Lewis type 2 antigen. H. ***pylori*** from patients with. Lewis(a+b-) expressed Lewis x more than y (1147 +/- 143 vs. 467 +/- 128 optical density units [ODU]; P = 0.006), isolates from patients with Lewis(a-b+) expressed Lewis x less than y (359 +/- 81 vs. 838 +/- 96 ODU; P = 0.0001), and isolates from Lewis(a-b-) patients expressed Lewis x and y approximately equally. ***Gastritis*** was unrelated to H. ***pylori*** Lewis expression. Conclusions: In mimicking host ***gastric*** epithelium, H. ***pylori*** cells not only express Lewis x and y, but the relative proportion of expression corresponds to the host Lewis phenotype, suggesting selection for host-adapted organisms.

L11 ANSWER 109 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 97:737204 SCISEARCH

GA The Genuine Article (R) Number: XY551

TI Helicobacter ***pylori*** infection in an Australian regional city: prevalence and risk factors

AU Peach H G (Reprint); Pearce D C; Farish S J

CS UNIV MELBOURNE, DEPT PUBL HLTH & COMMUNITY MED, BALLARAT HLTH SERV BASE HOSP, POB 577, BALLARAT, VIC 3353, AUSTRALIA (Reprint); UNIV MELBOURNE, EPIDEMIOL & BIOSTAT UNIT, DEPT PUBL HLTH & COMMUNITY MED, PARKVILLE, VIC 3052, AUSTRALIA

CYA AUSTRALIA

SO MEDICAL JOURNAL OF AUSTRALIA, (15 SEP 1997) Vol. 167, No. 6, pp. 310-313. Publisher: AUSTRALASIAN MED PUBL CO LTD, LEVEL 1, 76 BERRY ST, SYDNEY NSW 2000, AUSTRALIA.

ISSN: 0025-729X.

DT Article; Journal

FS LIFE; CLIN

LA English

REC Reference Count: 23

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

Objective: To investigate the prevalence of Helicobacter ***pylori*** infection and potential risk factors for infection in an adult Australian population.

Design: Cross-sectional study.

Setting: Ballarat, a major regional city in Victoria (population, 78 000; 92% born in Australia), November 1994 to July 1995.

Participants: 217 adults randomly selected from the electoral roll. Main outcome measures: H. ***pylori*** IgG ***antibody*** status by enzyme ***immunoassay***; amount of dental plaque; sociodemographic and other potential risk factors; odds ratios for risk factors determined by logistic regression analysis.

Results: Age-standardised prevalence of H. ***pylori*** infection was 30.6%. After adjustment for age, sex and socioeconomic index, positive H. ***pylori*** status was significantly associated with increasing number of tooth surfaces with a high plaque score (odds ratio [OR], 1.7; 95% confidence interval [CI], 1.1-2.7), increasing number of years in a job with public contact (OR, 1.7; 95% CI, 1.3-2.3), blood group B antigen (OR, 3.1 95% CI, 1.1-9.1), and having lived in a household with more than six members during childhood (OR, 2.5; 95% CI, 1.1-5.5). Negative H.

pylori status was significantly associated with increasing education, having ever lived on a farm, and having teeth scaled less than once a year.

Conclusions: H. ***pylori*** infection is common. Dental plaque may be a reservoir for H. ***pylori***, which is probably transmitted by person-to-person contact, and blood group B antigen may predispose to infection. Community education about effective oral hygiene and adoption of good hygiene practices by those with regular public contact may be important to prevent acquisition and transmission of H. ***pylori***.

- L11 ANSWER 110 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)
- AN 97:379884 SCISEARCH
- GA The Genuine Article (R) Number: WY184
- TI Evaluation of a rapid, new method for detecting serum IgG ***antibodies*** to Helicobacter ***pylori***
- AU Sharma T K; Young E L; Miller S; Cutler A F (Reprint)
- CS SINAI HOSP, GASTROENTEROL SECT, 6767 W OUTER DR, DETROIT, MI 48235 (Reprint), SINAI HOSP, GASTROENTEROL SECT, DETROIT, MI 48235 CYA USA
- SO CLINICAL CHEMISTRY, (MAY 1997) Vol. 43, No. 5, pp. 832-836. Publisher: AMER ASSOC CLINICAL CHEMISTRY, 2101 L STREET NW, SUITE 202, WASHINGTON, DC 20037-1526. ISSN: 0009-9147.

DT Article; Journal

FS LIFE

LA English

REC Reference Count: 26

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

There is an increased need for rapid, inexpensive tests to diagnose Helicobacter ***pylori*** infection. Our objective was to determine the performance characteristics of an immunochromatographic test (ICT) for detection of anti-H. ***pylori*** IgG ***antibodies*** . A commercially available ICT kit. (FlexSure(R) HP) was tested with a well-characterized cohort of banked sera as well as with fresh serum from randomly selected symptomatic patients. The ICT was evaluated with 107

stored sera and 96 prospective patients. The test correctly identified 65 of 68 H. ***pylori*** -infected and 37 of 39 noninfected stored sera and 54 of 57 infected and 30 of 39 noninfected patients. Sensitivity, specificity, and positive and negative predictive values were 96%, 95%, 97%, and 93% in stored serum and 95%, 77%, 86%, and 91% in fresh serum, respectively. We concluded that the ICT, reported at 4 min, is highly sensitive for detecting anti-H. ***pylori*** IgG ***antibodies*** in human serum. With a high negative predictive value, the test may be used to exclude H. ***pylori*** infection in symptomatic patients.

L11 ANSWER 111 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 97:137683 SCISEARCH

GA The Genuine Article (R) Number: WG312

TI Risk of infection with Helicobacter ***pylori*** and hepatitis A virus in different groups of hospital workers

AU Rudi J (Reprint); Toppe H; Marx N; Zuna I; Theilmann L; Stremmel W; Raedsch R

CS UNIV HEIDELBERG, DEPT MED, DIV GASTROENTEROL, BERGHEIMER STR 58, D-69115 HEIDELBERG, GERMANY (Reprint); ST JOSEF HOSP, DEPT MED, DIV GASTROENTEROL, WIESBADEN, GERMANY; GERMAN CANC RES CTR, RES PROGRAM RADIOL DIAGNOST & THERAPY, D-6900 HEIDELBERG, GERMANY

CYA GERMANY

SO AMERICAN JOURNAL OF GASTROENTEROLOGY, (FEB 1997) Vol. 92, No. 2, pp. 258-262

Publisher: WILLIAMS & WILKINS, 351 WEST CAMDEN ST, BALTIMORE, MD

21201-2436.

ISSN: 0002-9270.

DT Article; Journal

FS CLIN

LA English

REC Reference Count: 34

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

Objectives: The purpose of this study was to determine whether different staff groups in an acute care hospital are at increased risk of acquiring Helicobacter ***pylori*** and hepatitis A virus infection. Methods: We examined staff members of an acute care hospital for serum ***antibodies*** to H. ***pylori*** IgG (n = 457) and to hepatitis A virus (n = 434). The staff members were assigned to three groups: 1) nonmedical staff (n = 110), 2) medical and nursing staff (n = 272), and 3) medical and nursing staff working in a gastroenterology and endoscopy unit (n = 75). Serum ***antibodies*** were measured by validated enzyme ***immunoassays*** . A questionnaire inquiring about medical and professional history, history of upper GI pain and ulcer, as well as about the use of nonsteroidal antiinflammatory drugs or medication for GI complaints and smoking habits was completed by each person. Results: The seroprevalence of H. ***pylori*** was 35.5% in group I, 34.6% in group II, and 24.0% in group III (not significant). The seroprevalence of N. ***pylori*** ***antibodies*** increased with age (p < 0.001), and ***antibodies*** were present more frequently in women than in men (36.2) vs 25.4%, p < 0.05). After adjustment for age, duration of experience and the number of years working in the gastroenterology or endoscopy unit did not increase H. ***pylori*** seropositivity. No significant association was found between H. ***pylori*** seropositivity and history of upper GI pain, ulcers, use of nonsteroidal anti-inflammatory drugs or medication for Gf complaints, or tobacco use. The prevalence of

hepatitis A ***antibodies*** was similiar in the three groups (group I, 26.4%; II, 26.5%; III, 21.7%; not significant). Cross-tabulation showed that 67 subjects (15.4%) were seropositive for both H. ***pylori*** and hepatitis A (p < 0.001) and that 245 (56.5%) were negative for both. Seventy-seven (17.7.%) and 45 (10.4%) were seropositive for only H. ***pylori*** and for only hepatitis A, respectively. Conclusions: Occupational exposure to patients in an acute care hospital as well as to patients and to endoscopic procedures of a gastroenterology and endoscopy unit does not increase the rate of infection with H. ***pylori***. The significant correlation between the seroprevalences of H. ***pylori*** and hepatitis A ***antibodies*** suggests fecal-oral transmission of H. ***pylori***.

L11 ANSWER 112 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 97:39325 SCISEARCH

GA The Genuine Article (R) Number: WA283

TI Association between Helicobacter ***pylori*** infection and serum ***pepsinogen*** concentrations in gastroduodenal disease

AU Matsumoto K (Reprint); Konishi N; Ohshima M; Hiasa Y; Kimura E; Samori T

CS JAPAN CLIN LABS INC, DEV SECT, 5-16-26 MINAMISUITA, SUITA, OSAKA 564, JAPAN (Reprint), NARA MED UNIV, DEPT PATHOL 2, KASHIHARA, NARA 634, JAPAN CYA JAPAN

SO JOURNAL OF-CLINICAL PATHOLOGY, (DEC 1996) Vol. 49, No. 12, pp. 1005-1008. Publisher: BRITISH MED-JOURNAL PUBL GROUP, BRITISH MED ASSOC HOUSE, TAVISTOCK SQUARE, LONDON, ENGLAND WC1H 9JR. ISSN: 0021-9746.

DT Article; Journal

FS LIFE; CLIN

LA English

REC Reference Count: 37

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB Aim-To investigate the association between Helicobacter ***pylori***
infection and serum ***pepsinogen*** (PG)1 and 2 concentrations in
various gastroduodenal diseases.

Methods-Serum PG1 and 2 concentrations and ***antibodies*** to H
pylori were measured by enzyme linked immunosorbent assay (ELISA);

gastric mucosal pH was assessed and urease activity in biopsy tissue was determined. A comparison of the ELISA and urease test results permitted division of the cases into positive, false positive, false negative and negative categories for control, ***gastritis***, and ulcer groups.

Results-The ***gastric*** mucosal pH and serum PG2 in cases positive for H ***pylori*** were significantly increased in ulcer and ***gastritis*** cases compared with H ***pylori*** negative cases. Similar tendencies were observed for the false positive and false negative categories.

Conclusions-A positive ELISA reaction for ***antibodies*** and an increased serum PG2 concentration are reliable indicators of H ***pylori*** infection.

L11 ANSWER 113 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 96:183295 SCISEARCH

GA The Genuine Article (R) Number: TX796

TI QUANTITATIVE DETECTION OF SECRETORY IMMUNOGLOBULIN-A TO HELICOBACTER-***PYLORI*** IN ***GASTRIC*** -JUICE - ***ANTIBODY*** -CAPTURE

Va

ENZYME-LINKED-IMMUNOSORBENT-ASSAY

- AU HAYASHI S (Reprint); SUGIYAMA T; HISANO K; AWAKAWA T; KUROKAWA I; YACHI A; ISOGAI H; ISOGAI E; YOKOTA K; HIRAI Y; OGUMA K; FUJII N
- CS SAPPORO MED UNIV, SCH MED, DEPT MICROBIOL, S 1 W 17, SAPPORO, HOKKAIDO 060, JAPAN (Reprint); SAPPORO MED UNIV, SCH MED, DIV LAB DIAG, SAPPORO. HOKKAIDO 060, JAPAN; SAPPORO MED UNIV, SCH MED, DEPT INTERNAL MED, SECT 1, SAPPORO, HOKKAIDO 060, JAPAN; SAPPORO MED UNIV, SCH MED, DIV ANIM EXPERIMENTAT, SAPPORO, HOKKAIDO 060, JAPAN; HLTH SCI UNIV HOKKAIDO, SCH DENT, DEPT PREVENT DENT, ISHIKARI, HOKKAIDO, JAPAN; OKAYAMA UNIV, SCH MED, DEPT BACTERIOL, OKAYAMA 700, JAPAN

CYA JAPAN

SO JOURNAL OF CLINICAL LABORATORY ANALYSIS, (1996) Vol. 10, No. 2, pp. 74-77. ISSN: 0887-8013.

DT Article: Journal

FS LIFE

LA ENGLISH

REC Reference Count: 18

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

Helicobacter ***pylori*** is a major etiologic agent in gastroduodenal disorders. In this study, immunoglobulin A (IgA) ***antibodies*** to H. ***pylori*** were estimated in serum and ***gastric*** juice specimens from patients with ***gastritis*** and peptic ulcers using ***antibody*** capture enzyme-linked immunosorbent assays (ACELISAs). The ***antibody*** titers of the ACELISAs are independent of the ***antibody*** concentration and reflect the ratio of H. ***pylori*** -specific IgA to total IgA. The ratio is stable, although the ***antibody*** concentration fluctuates in ***gastric*** juice. Using the ACELISAs it was possible to evaluate quantitatively not only serum IgA (SR-IgA) ***antibodies*** but also secretory IgA (SC-IgA) ***antibodies*** in ***gastric*** juice. There were significant differences between the patients and control group in the SR-IgA and SC-IgA ACELISAs. Furthermore, the ACELISAs made it possible to compare between SR-IgA ***antibodies*** in serum and SC-IgA ***antibodies*** in ***gastric*** juice. In all patients, the ratios of H. ***pylori*** -specific IgA were higher in ***gastric*** juice than in serum. These results suggest that H. ***pylori*** SC-IgA ***antibodies*** are mainly produced by the local immune response in the ***gastric*** mucosa. Our studies indicate that ACELISA is well suited for the analysis of local immune

- L11 ANSWER 114 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)
- AN 95:687270 SCISEARCH
- GA The Genuine Article (R) Number: RX016

response in mucosa. (C) 1996 Wiley-Liss, Inc.

- TI HELICOBACTER- ***PYLORI*** INFECTION IN FINNISH CHILDREN AND ADOLESCENTS - A SEROLOGIC CROSS-SECTIONAL AND FOLLOW-UP-STUDY
- AU ASHORN M (Reprint); MAKI M; HALLSTROM M; UHARI M; AKERBLOM H K; VIIKARI J; MIETTINEN A
- CS UNIV TAMPERE, DEPT CLIN MED, POB 607, SF-33101 TAMPERE, FINLAND (Reprint): TAMPERE UNIV HOSP, DEPT CLIN MED, TAMPERE, FINLAND: UNIV OULU, DEPT PEDIAT, SF-90100 OULU, FINLAND; UNIV HELSINKI, CHILDRENS HOSP, DEPT PEDIAT 2, HELSINKI, FINLAND; TURKU UNIV, DEPT MED, TURKU, FINLAND CYA FINLAND
- SO SCANDINA VIAN JOURNAL OF GASTROENTEROLOGY, (SEP 1995) Vol. 30, No. 9, pd. 876-879.

ISSN: 0036-5521.
DT Article; Journal
FS LIFE; CLIN
LA ENGLISH

REC Reference Count: 18

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

Background: The purpose was to examine the epidemiology of Helicobacter ***pylori*** infection in Finnish children and adolescents. Methods: Blood samples taken from healthy subjects (n = 461) 3-18 years old were studied cross-sectionally for the presence of H. ***pylori*** ***antibodies*** . Additionally, blood samples drawn in 1980, 1983, 1986, and 1989 from 74 children born in 1977 were tested. Serum IgG-class ***antibodies*** to H. ***pylori*** were determined by an enzyme ***immunoassay*** Results: In the cross-sectional series the mean ***antibody*** levels and the percentage of seropositive children increased with age. The overall seroprevalence was 10.2%. During the follow-up period from 3 to 12 pars of age the seropositivity increased from 4.6% to 5.7%. On the basis of the seroconversions between 3 and 12 years of age the annual incidence of H. ***pylori*** infection was calculated to be only 0.3%. Conclusions: In children seropositivity for H. ***pylori*** of the IgG class is often a sign of an infection acquired in early childhood. It seems likely that the age-dependent increase in the seropositivity reflects cumulation of a chronic infection.

L11 ANSWER 115 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 95:549348 SCISEARCH

GA The Genuine Article (R) Number: RN498

TI HELICOBACTER- ***PYLORI*** SEROPOSITIVITY AMONG SWEDISH ADULTS WITH AND WITHOUT ABDOMINAL SYMPTOMS - A POPULATION-BASED EPIDEMIOLOGIC-STUDY

AU AGREUS L (Reprint); ENGSTRAND L, SVARDSUDD K, NYREN O, TIBBLIN G

CS VARDCENTRALEN, S-74221 OSTHAMMAR, SWEDEN (Reprint); UNIV UPPSALA, DEPT FAMILY MED, CLIN EPIDEMIOL UNIT, S-75105 UPPSALA, SWEDEN; UNIV UPPSALA, DEPT CLIN MICROBIOL, S-75105 UPPSALA, SWEDEN; UNIV UPPSALA HOSP, DEPT CANC EPIDEMIOL, UPPSALA, SWEDEN; PRIMARY HLTH CARE CTR, OSTHAMMAR, SWEDEN CYA SWEDEN

SO SCANDINAVIAN JOURNAL OF GASTROENTEROLOGY, (AUG 1995) Vol. 30, No. 8, pp. 752-757.

ISSN: 0085-5928.

DT Article; Journal

FS LIFE; CLIN

LA ENGLISH

REC Reference Count: 25

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB Background: The role of Helicobacter ***pylori*** in functional dyspepsia is unclear. The aim of this population-based study was to determine whether the prevalence of H. ***pylori*** infection is higher among people with dyspepsia or irritable bowel syndrome (IBS) than among symptomless persons after control for age, sex, and socioeconomic status. Methods: In a postal questionnaire we asked a representative sample (20-79 years; n = 1260) from a Swedish municipality about abdominal symptoms in the preceding 3 months. A randomly selected subsample, 50 with dyspepsia, 50 with IBS, and 50 symptomless, matched with regard to age, sex, and education, were tested for the presence of IgG ***antibodies*** to H. ***pylori***, using the HM-CAP ***immunoassay***. Results: Fifty-five persons (38%) were H. ***pylori*** -seropositive. The

seroprevalence among dyspeptics (33%) did not exceed that in healthy people (48%) or in those reporting IBS (33%). The prevalence increased with age and with lower social class, but the latter association disappeared when age was taken into account. Neither sex nor symptom intensity predicted Helicobacter seropositivity. Conclusion: Our data are incompatible with an important aetiologic role for H. ***pylori*** in functional dyspepsia.

L11 ANSWER 116 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 94:770438 SCISEARCH

GA The Genuine Article (R) Number: PV007

TI THE PREVALENCE OF HELICOBACTER- ***PYLORI*** POSITIVITY IN HUMAN IMMUNODEFICIENCY VIRUS-INFECTED CHILDREN

AU BLECKER U (Reprint); KEYMOLEN K; LANCIERS S; BAHWERE P; SOUAYAH H; LEVY J; VANDENPLAS Y

CS FREE UNIV BRUSSELS, ACAD CHILDRENS HOSP, DEPT PEDIAT GASTROENTEROL, LAABEEKLAAU 101, B-1090 BRUSSELS, BELGIUM (Reprint); FREE UNIV BRUSSELS, ACAD CHILDRENS HOSP, DEPT ANESTHESIOL, B-1090 BRUSSELS, BELGIUM; HOP UNIV ST PIERRE, DEPT PEDIAT, BRUSSELS, BELGIUM

CYA BELGIUM

SO JOURNAL OF PEDIATRIC GASTROENTEROLOGY AND NUTRITION, (NOV 1994) Vol. 19, No. 4, pp. 417-420.

ISSN: 0277-2116.

DT Article; Journal

FS LIFE; CLIN

LA ENGLISH

REC Reference Count: 13

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

To investigate the prevalence of Helicobacter ***pylori*** infection in pediatric patients infected with the human immunodeficiency virus, we sought to detect the presence of ***antibodies*** against this organism in 23 human immunodeficiency virus-infected children of central African ethnic origin by means of a second-generation enzyme-linked ***immunoassay*** (ELISA) test for the detection of immunoglobulin G (IgG) ***antibodies*** to Helicobacter ***pylori*** (Malakit Helicobacter ***pylori***, Biolab, Limal, Belgium). They were compared to an asymptomatic control population matched for age and ethnic origin. Blood samples were taken during routine blood analysis before the monthly administration of intravenous gamma-globulins in the human immunodeficiency virus-infected patients and during preoperative blood analysis in the control population. Despite the fact that most human immunodeficiency virus-infected patients had IgG ***antibodies*** against other frequently encountered pathogens, none of them had a positive serology for Helicobacter ***pylori***, compared to 10 of 52 patients (19.2%) in the control population. This difference is statistically significant (p = 0.01).

- L11 ANSWER 117 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)
- AN 94:743762 SCISEARCH
- GA The Genuine Article (R) Number: PT564
- TI SERUM ***ANTIBODY*** -RESPONSE TO THE SUPERFICIAL AND RELEASED COMPONENTS OF HELICOBACTER- ***PYLORI***
- AU BAZILLOU M; FENDRI C; CASTEL O; INGRAND P; FAUCHERE J L (Reprint)
- CS CHU LA MILETRIE, MICROBIOL LAB A, BP 577, F-86021 POITIERS, FRANCE (Reprint); CHU LA MILETRIE, MICROBIOL LAB A, F-86021 POITIERS, FRANCE; FAC

MED & PHARM POITIERS, DEPT PEDAG & INFORMAT MED, F-86031 POITIERS, FRANCE; HOSP LA RABTA, MICROBIOL LAB, TUNIS, TUNISIA

CYA FRANCE; TUNISIA

SO CLINICAL AND DIAGNOSTIC LABORATORY IMMUNOLOGY, (MAY 1994) Vol. 1, No. 3, pp. 310-317.

ISSN: 1071-412X. DT Article; Journal

FS CLIN

LA ENGLISH

REC Reference Count: 32

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

Superficial and released components were extracted from six selected Helicobacter ***pylori*** strains. The protein and antigenic profiles of these extracts were representative of the profiles found most frequently among the clinical strains and included major peptidic fractions at 19, 23.5, 57, 68, 76, 118, and 132 kDa and major antigens at 68, 57, and 23.5 kDa. Immuno-cross-reactions were seen vith a hyperimmune rabbit serum to Campylobacter fetus but not with sera to Campylobacter jejuni or Salmonella spp. An antigenic preparation was obtained by pooling equivalent quantities of each extract, and the antigenic preparation, vas used to study the ***antibody*** responses of sera from 65 French patients and 127 Tunisian patients. By enzyme-linked immunosorbent assay, we observed that the sera from French and Tunisian patients clustered into two populations, defined as ***antibody*** positive (72 patients) and ***antibody*** negative (120 patients). The ***antibody*** -positive patients were more frequently infected with H. ***pylori*** (P < 0.01) and were more frequently affected with ***gastritis**** (P = 0.05). However, no correlation between ***antibody*** levels and clinical signs of dyspepsia was noticed. The proportions of ***antibody*** -positive patients were similar in France and Tunisia. ***Antibody*** -positive and ***antibody*** -negative sera were studied by Western blot (immunoblot) analysis. The ***antibody*** -positive sera revealed an average of 7.7 antigenic bands, whereas the ***antibody*** -negative sera revealed an average of 2.4 antigenic bands (P < 0.01). The antigens between 15 and 40 kDa and greater than 66 kDa were specifically recognized by the ***antibody*** -positive sera, although in this molecular size range the ***antibody*** profiles of these sera exhibited a fairly high degree of diversity. We conclude that the superficial and released components from H. ***pylori*** contain a variety of bacterial immunogens and mag be useful in antigenic preparations for the serodiagnosis of H. ***pylori*** infections. Moreover, a group of antigens in combination appears to be useful for discriminating ***antibody*** -positive and ***antibody*** -negative patients.

L11 ANSWER 118 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 93:634261 SCISEARCH

GA The Genuine Article (R) Number: MB444

TI HELICOBACTER- ***PYLORI*** INFECTION IN VARIOUS GROUPS OF GENERAL-HOSPITAL INPATIENTS AND DONORS

AU KALININ A V (Reprint); SPESIVTSEV V N; SKVORTSOV S V; LYTSAR B N

CS NN BURDENKO CENT MIL CLIN HOSP, MOSCOW, RUSSIA (Reprint)

CYA RUSSIA

SO KLINICHESKAYA MEDITSINA, (1993) Vol. 71, No. 3, pp. 38-39. ISSN: 0023-2149.

DT Article; Journal

FS CLIN

LA Russian

REC Reference Count: 8

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

Incidence of Helicobacter ***pylori*** (HP) infection in inpatients of a general hospital and donors was evaluated immunologically: the serum was examined for IgG ***antibodies*** to HP using enzyme ***immunoassay*** . Of 354 examinees 89 had duodenal ulcer (DU), 101 were healthy donors. HP infection was found prevalent in all the groups of the inpatients. Cases of HP were more numerous in the gastroenterological department as compared to other departments (81.6% against 39.6%. respectively). Among patients of the gastroenterological department DU patients were most frequent HP carriers (93.2%), this being indicative of a close correlation between DU and HP revealed previously by other diagnosic methods. Rare occurrence of HP in normal subjects (6-11%) reported by many authors was not confirmed, as HP was detected in 33% of the donors. Age-related analysis points to early onset of HP infection (25-26% incidence in 18-20-year-olds), its peak in middle-aged and presenile patients (78-80% in 50-69-year-olds) and lower occurrence in senile patients (25% at the age over 70).

L11 ANSWER 119 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 93:629733 SCISEARCH

GA The Genuine Article (R) Number: MB612

TI PERFORMANCE OF HELICOBACTER- ***PYLORI*** ACID EXTRACT AND UREASE ENZYME-LINKED IMMUNOSORBENT ASSAYS IN RELATION TO C-14 UREA BREATH TEST

AU VONWULFFEN H (Reprint); GATERMANN S; WINDLER E; GABBE E; HEINRICH H C

CS UNIV KRANKENHAUS EPPENDORF, INST MED MIKROBIOL & IMMUNOL, MARTINSTR 52, D-20246 HAMBURG, GERMANY (Reprint); MED UNIV LUBECK, INST MED MIKROBIOL, D-23562 LUBECK, GERMANY; UNIV KRANKENHAUS EPPENDORF, MED KERNKLIN, D-20246 HAMBURG, GERMANY; UNIV KRANKENHAUS EPPENDORF, MED BIOCHEM ABT, D-20246 HAMBURG, GERMANY

CYA GERMANY

SO ZENTRALBLATT FUR BAKTERIOLOGIE-INTERNATIONAL JOURNAL OF MEDICAL MICROBIOLOGY VIROLOGY PARASITOLOGY AND INFECTIOUS DISEASES, (SEP 1993) Vol. 280, No. 1-2, pp. 203-213.

ISSN: 0934-8840.

DT Article: Journal

FS LIFE

LA ENGLISH

REC Reference Count: 22

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

The C-14-urea breath test has been shown to be a reliable non-invasive method to detect the presence or absence of H. ***pylori*** infection. Alternatively, a number of techniques have been devised to detect circulating ***antibodies*** against H. ***pylori*** in serum, the most commonly used being enzyme-linked immunosorbent assays (ELISA). In the present study we compared the value of two ELISA antigen preparations, an acid glycine extract and a urease preparation, in relation to the results achieved in a C-14-urea breath test. Seventy-five gastroenterology outpatients were screened for the presence of H. ***pylori*** infection using the urea breath test. At the same time serum specimens were obtained. Thirty-seven patients had a positive breath test, i.e. they expired more than 2% of the oral C-14 test dose within 60 min. Using the breath test as reference, sensitivity and specificity for the acid extract

were 89.2% and 84.2% respectively, and for the urease ELISA 81.1% and 89.5%. Agreement between the two ELISAs was found in 82.7%, overall agreement between all three tests was observed in 77.3%. All three tests were found to be useful for monitoring therapy directed against H.

pylori

L11 ANSWER 120 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 93:320262 SCISEARCH

GA The Genuine Article (R) Number: LC040

TI USEFULNESS OF SEVERAL COMMERCIAL ENZYME-LINKED ***IMMUNOASSAYS*** FOR DETECTION OF HELICOBACTER- ***PYLORI*** INFECTION IN CLINICAL MEDICINE

AU LOFFELD R J L F (Reprint); VRIESE W T J; STOBBERINGH E E

CS ZIEKENHUIS DE HEEL, DEPT INTERNAL MED, POB 210, 1500 EE ZAANDAM, NETHERLANDS (Reprint)

CYA NETHERLANDS

SO EUROPEAN JOURNAL OF GASTROENTEROLOGY & HEPATOLOGY, (MAY 1993) Vol. 5, No. 5, pp. 333-337.

ISSN: 0954-691X.

DT Article; Journal

FS CLIN

LA ENGLISH

REC No References Keyed

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB Objective: To study six commercial enzyme-linked immunosorbent assays (ELISA) for detection of Helicobacter ***pylori*** immunoglobulin (Ig)G ***antibodies***.

Design: A comparison of different ELISAs in patients with known H. ***pylori*** status.

Patients: Patients who underwent endoscopy during which biopsy specimens were taken for detection of H. ***pylori*** via standard histologic and microbiologic techniques.

Results: Positive and negative predictive values for the different ELISAs are comparable, but major inter- and intra-assay variability was present.

Conclusion: Commercial assays should be tested with a local reference population in order to obtain the optimal cut-off value via a receiver operating characteristics curve. The use of the cut-off values provided by the manufacturers introduces difficulties with interpretation and is not valid.

L11 ANSWER 121 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 92:557059 SCISEARCH

GA The Genuine Article (R) Number: JN651

TI EFFECT OF ANTIMICROBIAL THERAPY ON THE SPECIFIC SEROLOGICAL RESPONSE TO HELICOBACTER- ***PYLORI*** INFECTION

AU GLUPCZYNSKI Y (Reprint); BURETTE A; GOOSSENS H; DEPREZ C; BUTZLER J P

CS BRUGMANN UNIV HOSP, DEPT CLIN MICROBIOL, 4 PL A VAN GEHUCHTEN, B-1020 BRUSSELS, BELGIUM (Reprint); NOUVELLE CLIN BASILIQUE, GASTROENTEROL UNIT, B-1080 BRUSSELS, BELGIUM; ST PIETERS HOSP, B-1000 BRUSSELS, BELGIUM; BRUGMANN UNIV HOSP, DEPT PATHOL, B-1020 BRUSSELS, BELGIUM; WHO, COLLABORATING CTR ENTER CAMPYLOBACTER, B-1000 BRUSSELS, BELGIUM CYA BELGIUM

SO EUROPEAN JOURNAL OF CLINICAL MICROBIOLOGY & INFECTIOUS DISEASES, (JUL 1992) Vol. 11, No. 7, pp. 583-588. ISSN: 0934-9723.

DT Article; Journal FS LIFE

LA ENGLISH

REC Reference Count: 33

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

The systemic immune response to Helicobacter ***pylori*** was studied in 247 infected adult patients before antimicrobial therapy and at different intervals following therapy. Endoscopy with simultaneous collection of biopsies was performed in all patients immediately before treatment, 4 to 6 weeks after the end of therapy and 6 to 12 months later. A C-14-urea breath test was performed 3 to 6 months after the end of treatment. Biopsy specimens were cultured and examined histologically using Giemsa stain. Sera were tested for Helicobacter ***pylori*** IgG ***antibodies*** with a commercial enzyme ***immunoassay*** using species-specific antigens. Overall, Helicobacter ***pylori*** was eradicated in 120 patients while the other 1127 remained infected with the organism. The follow-up period ranged from 4 weeks to 33 months (mean 10.2 months). Pretreatment IgG levels did not differ significantly between the two groups of patients. Six weeks after the end of treatment a slight but definite decrease in the IgG ***antibody*** levels was seen irrespective of treatment success. In the 127 patients who remained Helicobacter ***pylori*** -positive, the level of IgG ***antibodies*** remained stable or increased with time. A continuous fall in ***antibody*** levels was observed following bacterial eradication in the other 120 patients, but the difference in ***antibody*** levels between treatment responders and nonresponders became significant only more than six months after the end of treatment (p = 0.001). Serological testing may be useful for monitoring the outcome of long-term treatment of Helicobacter ***pylori*** infection and obviate

L11 ANSWER 122 OF 184 SCISEARCH COPYRIGHT 2001 ISI (R)

AN 92:140270 SCISEARCH

the need for endoscopy.

GA The Genuine Article (R) Number: HG098

TI EFFECT OF ALCOHOL-CONSUMPTION ON THE RISK OF HELICOBACTER- ***PYLORI***
INFECTION

AU HOOKNIKANNE J (Reprint)

CS UNIV HELSINKI, DEPT BACTERIOL & IMMUNOL, HAARTMANINKATU 3, SF-00290 HELSINKI 29, FINLAND (Reprint)

CYA FINLAND

SO DIGESTION, (OCT 1991) Vol. 50, No. 2, pp. 92-98. ISSN: 0012-2823.

DT Article: Journal

FS LIFE; CLIN

LA ENGLISH

REC Reference Count: 30

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB To investigate the effect of alcohol consumption on the risk of Helicobacter ***pylori*** infection, standardized questionnaires on drinking habits were used to interview 451 patients, whose H.

****pylori*** status was determined both by culture and serology.

Reported alcohol consumption did not increase the risk of H.

pylori infection (a 1.0 odds ratio, CI95 0.6-1.6). However, when the patients were divided into two age-groups, those under 35 years who reported to use alcohol seemed to have a slightly higher risk of H.

pylori infection (a 3.3 odds ratio CI95 0.9-12.2) compared to those over 35 years (a 1.0 odds ratio, CI95 0.5-2.2). This phenomenon did not reach statistical significance. The type of alcohol consumed did not affect the age-adjusted risk of H. ***pylori*** infection. If pathologically defined chronic ***gastritis*** was found, the risk for H. ***pylori*** was high (a 26.7 odds ratio, CI95 12.1-59.0, for those under 35 years, and a 12.8 odds ratio, CI95 6.7-24.3, for those over 35 years of age).

L11 ANSWER 123 OF 184 USPATFULL

AN 2001:33009 USPATFULL

TI Human myosin heavy chain-like proteins and method of detecting nucleic acid encoding said proteins

IN Bandman, Olga, Mountain View, CA, United States
 Yue, Henry, Sunnyvale, CA, United States
 Corley, Neil C., Mountain Vew, CA, United States
 Shah, Purvi, Sunnyvale, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 6197512 20010306

AI US 1998-216619 19981217 (9)

RLI Division of Ser. No. US 1997-966318, filed on 7 Nov 1997, now patented, Pat. No. US 6001593

DT Utility

EXNAM Primary Examiner: Carlson, Karen Cochrane

LREP Incyte Pharmaceuticals, Inc

CLMN Number of Claims: 5

ECL Exemplary Claim: 1

DRWN 13 Drawing Figure(s); 13 Drawing Page(s)

LN.CNT 2334

AB The invention provides human myosin heavy chain-like proteins (MHCP) and polynucleotides which identify and encode MHCP. The invention also provides expression vectors, host cells, ***antibodies***, agonists, and antagonists. The invention also provides methods for treating disorders associated with expression of MHCP.

L11 ANSWER 124 OF 184 USPATFULL

AN 2000:174620 USPATFULL

TI Histidine kinase

IN Biswas, Sanjoy, Paoli, PA, United States
 Throup, John P, Royersford, PA, United States
 Wallis, Nicola G, Wayne, PA, United States
 Zalacain, Magdalena, West Chester, PA, United States

PA SmithKline Beecham Corporation, Philadelphia, PA, United States (U.S. corporation)

SmithKline Beecham p.l.c., United Kingdom (non-U.S. corporation)

PI US 6165992 20001226

AI US 1998-81689 19980520 (9)

PRAI US 1997-48347 19970530 (60)

DT Utility

EXNAM Primary Examiner: Priebe, Scott D.

LREP Gimmi, Edward R.; Deibert, Thomas S.; King, William T.

CLMN Number of Claims: 30

ECL Exemplary Claim: 1,16

DRWN No Drawings

LN.CNT 2877

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides Histidine kinase polypeptides and polynucleotides encoding Histidine kinase polypeptides and methods for producing such polypeptides by recombinant techniques. Also provided are methods for utilizing Histidine kinase polypeptides to screen for antibacterial compounds.

L11 ANSWER 125 OF 184 USPATFULL

AN 2000:174365 USPATFULL

TI. Method of detecting bacterial infection

IN Fawcett, Paul Thomas, Rising Sun, MD, United States

PA The Nemours Foundation, Wilmington, DE, United States (U.S. corporation)

PI US 6165736 20001226

AI US 1998-123231 19980728 (9)

DT Utility

EXNAM Primary Examiner: Bui, Phuong T.

LREP McGuireWoods, LLP CLMN Number of Claims: 18 ECL Exemplary Claim: 1

DRWN 1 Drawing Figure(s); 1 Drawing Page(s)

LN.CNT 1393

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A method of developing sensitive and discriminatory diagnostic procedures for detecting active bacterial infection in animals, especially humans, basically involves partially digesting the genomic DNA of the infecting bacterial pathogen into a generally large number of ideally random fragments and finding proteins encoded by those fragments which evoke a discriminating response to specimens from viably infected animals. Cloning techniques are used to cause the genes of the multitude of DNA fragments to produce proteins. Groups of proteins encoded by the genes of each fragment are separately tested for the ability to generate an immune response in certain specimens from animals known to have "viable infection", "convalescent infection" and "naive status" with respect to infection by the infecting bacterial pathogen. The protein groups which evoke positive immune responses to viably infected but no immune response to naive specimens are identified as "selectively responsive proteins". Similarly, selectively responsive proteins which are found to evoke no immune response from convalescent specimens are identified as "discriminatingly responsive proteins". These selectively and discriminatingly responsive protein groups can be cloned in magnitude and used to test unknown patients for status of infection.

The method is amenable for developing tests based upon non-invasively obtained specimens, such as peripherally-obtained blood samples. Moreover, rigorous mapping of the pathogen genome is not prerequisite for carrying out the development method. Consequently, the development method can be used to obtain diagnostic procedures particularly suitable for generating individually inexpensive bacterial infection assays capable for screening large scale patient populations.

L11 ANSWER 126 OF 184 USPATFULL

AN 2000:160787 USPATFULL

TI TagA gene and methods for detecting predisposition to peptic ulceration and ***gastric*** carcinoma

IN Cover, Timothy L., Nashville, TN, United States
 Blaser, Martin J., Nashville, TN, United States
 Kleanthous, Harry, Cambridge, MA, United States
 Tummuru, Murali K. R., Nashville, TN, United States

PA Vanderbilt University, Nashville, TN, United States (U.S. corporation)

PI US 6153390 20001128

AI US 1999-259437 19990301 (9)

RLI Continuation of Ser. No. US 1998-34306, filed on 2 Mar 1998, now patented, Pat. No. US 5876943 which is a division of Ser. No. US 1994-316397, filed on 30 Sep 1994, now patented, Pat. No. US 5733740 which is a continuation-in-part of Ser. No. US 1993-53614, filed on 26 Apr 1993, now patented, Pat. No. US 5403924 which is a continuation-in-part of Ser. No. US 1992-959940, filed on 13 Oct 1992, now abandoned

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Needle & Rosenberg, P.C.

CLMN Number of Claims: 4

ECL Exemplary Claim: 1

DRWN 4 Drawing Figure(s); 4 Drawing Page(s)

LN.CNT 2842

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides an isolated nucleic acid encoding an approximately 120-128 kilodalton antigen of Helicobacter ***pylori***, or an antigenic fragment thereof, wherein the antigen is associated with peptic ulceration. The present invention also provides methods of detecting the presence of a Helicobacter ***pylori*** strain possessing the 120-128 kilodalton antigen in a subject, comprising the steps of contacting an ***antibody*** -containing sample from the subject with a detectable amount of the tagA antigen or antigenic polypeptide of the present invention and detecting the binding of the antigen or fragment and the ***antibody***. The detection of a strain expressing the TagA antigen is an indication of predisposition to peptic ulceration and ***gastric*** carcinoma. A mutant H.

pylori not expressing a functional TagA antigen is also provided.

L11 ANSWER 127 OF 184 USPATFULL

AN 2000:149964 USPATFULL

TI Human lysophospholipase

IN Hillman, Jennifer L., Mountain View, CA, United States Shah, Purvi, Sunnyvale, CA, United States Corley, Neil C., Mountain View, CA, United States Murry, Lynn E., Portola Valley, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 6143544 20001107

AI US 1997-878862 19970619 (8)

DT Utility

EXNAM Primary Examiner: Hendricks, Keith D.; Assistant Examiner: Mayhew, Bradley S.

LREP Billings, Lucy J.; Price, Leanne C. Incyte Pharmaceuticals, Inc.

CLMN Number of Claims: 9

ECL Exemplary Claim: 1

DRWN 5 Drawing Figure(s); 5 Drawing Page(s)

LN.CNT 2199

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides a new human lysophospholipase (JHLP) and polynucleotides which identify and encode IHLP. The invention also provides expression vectors, host cells, agonists, ***antibodies*** and antagonists. The invention also provides methods for treating inflammation and disorders associated with expression of IHLP.

L11 ANSWER 128 OF 184 USPATFULL

AN 2000:145883 USPATFULL

TI ATP-dependent RNA helicase protein

IN Bandman, Olga, Mountain View, CA, United States Guegler, Karl J., Menlo Park, CA, United States Corley, Neil C., Mountain View, CA, United States Shah, Purvi, Sunnyvale, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 6139837 20001031

AI US 1998-149934 19980909 (9)

RLI Division of Ser. No. US 1997-892256, filed on 11 Jul 1997, now patented, Pat. No. US 5888792

DT Utility

EXNAM Primary Examiner: Slobodyansky, Elizabeth

LREP Incyte Pharmaceuticals, Inc.

CLMN Number of Claims: 4

ECL Exemplary Claim: 1

DRWN 14 Drawing Figure(s); 14 Drawing Page(s)

LN.CNT 2351

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides a human ATP-dependent RNA helicase (ADRH-1) and polynucleotides which identify and encode ADRH-1. The invention also provides expression vectors, host cells, agonists, ***antibodies*** and antagonists. The invention also provides methods for treating disorders associated with expression of ADRH-1.

L11 ANSWER 129 OF 184 USPATFULL

AN 2000:142121 USPATFULL

TI Microbiological media for isolation and indentification of enteric pathogens such as E. coli and salmonella

IN Bochner, Barry, Alameda, CA, United States

PA Biolog, Inc., CA, United States (U.S. corporation)

PI US 6136554 20001024

AI US 1997-819452 19970317 (8)

RLI Continuation of Ser. No. US 1995-484960, filed on 7 Jun 1995

DT Utility

EXNAM Primary Examiner: Marx, Irene

LREP Medlen & Carroll, LLP

CLMN Number of Claims: 21

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 3394

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention is directed to methods and media for the growth, enrichment, isolation, and presumptive identification of enteric

pathogens such as E. coli 0157:H7 and Salmonella. In particular, the organisms commonly associated with gastrointestinal infections of humans and other animals are distinguished based on their growth, colonial morphology and color. The present invention is also directed to methods and media for the growth, enrichment, isolation and presumptive identification of enteric pathogens such as E. coli 0157:H7 and Salmonella isolated from food, water, dairy, and environmental samples.

L11 ANSWER 130 OF 184 USPATFULL

AN 2000:134726 USPATFULL

TI Helicobacter ***pylori*** proteins useful for vaccines and diagnostics

IN Covacci, Antonello, Vc.Provenzano, 8, 53100, Siena, Italy
 Bugnoli, Massimo, V. del Pozzo, 38, 53035, Monteriggioni, Italy
 Telford, John, Via Sambuco, 43, 53010, Monteriggioni, Italy
 Macchia, Giovanni, Via Monte Velino 57, 67051 Avezzano (AQ), Italy
 Rappuoli, Rino, Via Calamandrei, 39, 53010 Quercegrossa (SI), Italy

PI US 6130059 20001010

AI US 1995-466662 19950606 (8)

RLI Division of Ser. No. US 1994-256848, filed on 21 Oct 1994, now abandoned which is a continuation of Ser. No. WO 1993-EP472, filed on 2 Mar 1993

PRAI IT 1992-FI52 19920302

WO 1993-EP158 19930125

DT Utility

EXNAM Primary Examiner: Bui, Phuong T.

LREP Harbin, Alisa A.; Paintin, Francis A.; Blackburn, Robert P.

CLMN Number of Claims: 6 ECL Exemplary Claim: 1

DRWN 14 Drawing Figure(s); 14 Drawing Page(s)

LN.CNT 2756

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Helicobacter ***pylori*** is known to cause or be a cofactor in type B ***gastritis***, peptic ulcers, and ***gastric*** tumors. In both developed and developing countries, a high percentage of people are infected with this bacterium. The present invention relates generally to certain H. ***pylori*** proteins, to the genes which express these proteins, and to the use of these proteins for diagnostic and vaccine applications. Specifically, molecular cloning, nucteotide, and amino acid sequences for the H. ***pylori*** cytotoxin (CT), the "Cytotoxin Associated Immunodominant" (CAI) antigen, and the heat shock protein (hsp60) are described herein.

L11 ANSWER 131 OF 184 USPATFULL

AN 2000:113750 USPATFULL

TI F.sub.0 ATP synthase subunit

IN Hillman, Jennifer L., Mountain View, CA, United States Goli, Surya K., Sunnyvale, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 6110722 20000829

AI US 1998-66049 19980424 (9)

RLI Division of Ser. No. US 1997-815177, filed on 11 Mar 1997, now patented, Pat. No. US 5786150

DT Utility

EXNAM Primary Examiner: Patterson, Jr., Charles L.

LREP Hamlet-Cox, DianaIncyte Pharmaceuticals, Inc. CLMN Number of Claims: 10

ECL Exemplary Claim: 1

DRWN 4 Drawing Figure(s); 3 Drawing Page(s)

LN.CNT 1951

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides a human ATP synthase subunit (ASYS) and polynucleotides which encode ASYS. The invention also provides expression vectors, host cells, agonists, antisense molecules, ****antibodies****, or antagonists. The invention also provides methods for producing ASYS and for treating disorders associated with expression of ASYS.

L11 ANSWER 132 OF 184 USPATFULL

AN 2000:109968 USPATFULL

TI iceA gene and related methods

Miller, Geraldine G., Franklin, TN, United States
 Peek, Jr., Richard M., Nashville, TN, United States
 Thompson, Stuart A., Whites Creek, TN, United States
 Blaser, Martin J., Nashville, TN, United States

PA Vanderbilt University, Nashville, TN, United States (U.S. corporation)

PI US 6107464 20000822

AI US 1999-413140 19991006 (9)

RLI Division of Ser. No. US 1998-60584, filed on 15 Apr 1998, now patented, Pat. No. US 6004354 which is a division of Ser. No. US 1996-650528, filed on 20 May 1996, now patented, Pat. No. US 5780278, issued on 14 Jul 1998

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Needle & Rosenberg, P.C.

CLMN Number of Claims: 6

ECL Exemplary Claim: 1

DRWN 8 Drawing Figure(s); 7 Drawing Page(s)

LN.CNT 2188

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A purified IceA protein of Helicobacter ***pylori*** is provided. The protein is expressed as either an IceA 1 or an IceA 2 variant. A purified polypeptide fragment of the IceA protein is also provided. An antigenic fragment of IceA is provided. An isolated nucleic acid that encodes an IceA protein of H. ***pylori*** is provided. A nucleic acid that encodes an IceA 1 variant and a nucleic acid that encodes an IceA 2 variant is also provided. Fragments of the iceA gene are provided. A method of detecting the presence of an ***antibody*** against H. ***pylori*** in a sample is provided. The method comprises the following steps: a) contacting the sample with a purified IceA protein of H. ***pylori*** or a H. ***pylori*** -specific fragment thereof; and b) detecting the binding of the ***antibody*** in the sample to the protein or fragment, the detection of binding indicating the presence in the sample of ***antibodies*** against H. ***pylori*** . A method of detecting the presence of an ***antibody*** against an ulcerative Helicobacter ***pylori*** strain in a sample is also provided.

AN 2000:94860 USPATFULL

TI Human lysophospholipase

IN Hillman, Jennifer L., Mountain View, CA, United States Shah, Purvi, Sunnyvale, CA, United States Murry, Lynn E., Portola Valley, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 6093561 20000725

AI US 1998-216386 19981218 (9)

RLI Division of Ser. No. US 1998-22940, filed on 12 Feb 1998 which is a continuation-in-part of Ser. No. US 1997-844120, filed on 29 Apr 1997, now patented, Pat. No. US 5858756

DT Utility

EXNAM Primary Examiner: Achutamurthy, Ponnathapu; Assistant Examiner: Mayhew, Bradley S.

LREP Muenzen, Colette C.; Mohan-Peterson, SheelaIncyte Pharmaceuticals, Inc.

CLMN Number of Claims: 2

ECL Exemplary Claim: 1

DRWN 9 Drawing Figure(s); 9 Drawing Page(s)

LN.CNT 2310

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides a human lysophospholipase (NHLP) and polynucleotides which identify and encode NHLP. The invention also provides expression vectors, host cells, ***antibodies***, agonists, and antagonists. The invention also provides methods for treating or preventing disorders associated with expression of NHLP.

L11 ANSWER 134 OF 184 USPATFULL

AN 2000:91766 USPATFULL

TI Helicobacter ***pylori*** proteins useful for vaccines and diagnostics

IN Covacci, Antonello, Siena, ItalyBugnoli, Massimo, Monteriggioni, Italy

Telford, John, Monteriggioni, Italy

Macchia, Giovanni, Avezzano, Italy

Rappuoli, Rino, Quercegrossa, Italy

PA Chiron S.p.A., Siena, Italy (non-U.S. corporation)

PI US 6090611 20000718

AI US 1995-471491 19950606 (8)

RLI Division of Ser. No. US 256848

PRAI IT 1992-FI52 19920302

WO 1993-EP158 19930125

DT Utility

EXNAM Primary Examiner: Bui, Phoung T.

LREP Paintin, Francis, Harbin, Alisa A., Blackburn, Robert P.

CLMN Number of Claims: 10

ECL Exemplary Claim: 1

DRWN 14 Drawing Figure(s); 14 Drawing Page(s)

LN.CNT 2776

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A cytotoxin associated immunodominant antigen and the nucleic acid encoding the antigen from Helicobacter ***pylori*** are described. This antigen was identified from the cytotoxin positive CCUG 17874 Helicobacter ***pylori*** strain, and both the antigen and the DNA encoding it have been sequenced. The antigen is a hydrophilic,

surface-exposed protein having a molecular weight of 120-132 kDa. The nucleic acid encoding the antigen may be incorporated into a vector for transformation of host cells for expression of the antigen. Both the DNA and the antigen can be used in assays for detection of disease or infection by Helicobacter ***pylori***, and may find use in treating and preventing infection by Helicobacter ***pylori*** and the diseases associated with such infection.

L11 ANSWER 135 OF 184 USPATFULL

AN 2000:80564 USPATFULL

TI Helicobacter catalase nucleotide sequences, their production and use

IN Sugiyama, Tosiro, Hokkaido, Japan Kawabata, Tomohisa, Osaka, Japan Hirayasu, Kazunari, Osaka, Japan

Tanaka, Takumi, Hyogo, Japan

PA Wako Pure Chemical Industries, Ltd., Osaka, Japan (non-U.S. corporation)

PI US 6080556 20000627

AI US 1996-657868 19960531 (8)

PRAI JP 1995-136564 19950602

JP 1996-83512 19960405

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny

LREP Conlin, David G.; Neuner, George W.Dike, Bronstein, Roberts & Cushman LLP

CLMN Number of Claims: 7

ECL Exemplary Claim: 1

DRWN 11 Drawing Figure(s); 6 Drawing Page(s)

LN.CNT 1491

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Disclosed are amino acid sequences of polypeptides reacting with
antibodies to Helicobacter ***pylori*** (HP), DNAs coding
therefor, vectors containing said DNAs, transformants containing said
vectors, a method for preparing said polypeptides by cultivating said
transformants, and anti-HP ***antibody*** assaying reagents and HP
gene detecting reagents comprising said polypeptides, thereby enabling
specific, quantitative inspection of HP.

L11 ANSWER 136 OF 184 USPATFULL

AN 2000:77222 USPATFULL

TI Helicobacter ***pylori*** proteins useful for vaccines and diagnostics

IN Covacci, Antonello, Siena, Italy

Bugnoli, Massimo, Monteriggioni, Italy

Telford, John, Monteriggioni, Italy

Macchia, Giovanni, Avezzano, Italy

Rappuoli, Rino, Quercegrossa, Italy

PA Chiron Corporation, Emeryville, CA, United States (U.S. corporation)

PI US 6077706 20000620

AI US 1995-470260 19950606 (8)

RLI Division of Ser. No. US 256848

PRAI IT 1992-FI52 19920302

WO 1993-EP158 19930125

DT Utility

EXNAM Primary Examiner: Stucker, Jeffrey; Assistant Examiner: Bui, Phuong T.

LREP Paintin, Francis A.; Harbin, Alisa A.; Blackburn, Robert P.

CLMN Number of Claims: 4

ECL Exemplary Claim: 1

DRWN 14 Drawing Figure(s); 14 Drawing Page(s)

LN.CNT 2677

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Helicobacter ***pylori*** known to cause or be a cofactor in type B ***gastritis***, peptic ulcers, and ***gastric*** tumors. In both developed and developing countries, a high percentage of people are infected with this bacterium. The present invention relates generally to certain H. ***pylori*** proteins, to the genes which express these proteins, and to the use of these proteins for diagnostic and vaccine applications. Specifically, molecular cloning, nucleotide, and amino add sequences for the H. ***pylori*** cytotoxin (CT), the "Cytotoxin Associated Immunodominant" (CAI) antigen, and the heat shock protein (hsp60) are described herein.

L11 ANSWER 137 OF 184 USPATFULL

AN 2000:67576 USPATFULL

TI In vitro test for Helicobacter ***pylori***

IN Cripps, Allan, Curtin, Australia

Witt, Campbell, Bicton, Australia

Clancy, Robert Llewellyn, Newcastle, Australia

Stiel, Daniel, East Lindfield, Australia

PA Provalis UK Limited, Deeside, United Kingdom (non-U.S. corporation)

PI US 6068985 20000530

WO 9322682 19931111

AI US 1995-325264 19950426 (8)

WO 1993-GB894 19930429

19950426 PCT 371 date

19950426 PCT 102(e) date

PRAI CA 1992-2067603 19920429

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Foley & Lardner

CLMN Number of Claims: 20

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 766

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Contemporary infection by Helicobacter ***pylori*** in a human or other mammal can be detected by detecting H. ***pylori*** -specific IgG ***antibody*** in saliva, or other mucous secretion, with an antigen preparation from H. ***pylori***. Diagnosis depends on detection of the antigen- ***antibody*** complex. For improved reliability, the antigen preparation comprises H. ***pylori*** -derived components of about 265 kDa and about 340 kDa and is substantially free of an H. ***pylori*** -derived component of about 440 kDa. The antigen preparation may be immobilised onto a solid support such as a nitrocellulose strip.

L11 ANSWER 138 OF 184 USPATFULL

AN 2000:57536 USPATFULL

TI Compositions and methods relating to drug discovery and detection and

treatment of gastrointestinal diseases

IN Corthesy-Theulaz, Irene, Epalinges, Switzerland

PA Kieta Holding SA, St-Prex, Switzerland (non-U.S. corporation)

PI US 6060241 20000509

AI US 1997-834776 19970403 (8)

PRAI US 1996-14906 19960405 (60)

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Cooley Godward LLP

CLMN Number of Claims: 12

ECL Exemplary Claim: 1

DRWN 13 Drawing Figure(s); 10 Drawing Page(s)

LN.CNT 2585

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A poly-3-hydroxybutyrate metabolic pathway essential for Helicobacter ***pylori*** survival in a host is provided. A novel Helicobacter ***pylori*** Coenzyme A transferase (Hp CoA-t), thiolase and PHB synthase as well as methods for their preparation and use are provided. Hp CoA-t and thiolase polynucleotides and proteins are provided as well as detection and preparative methods using such molecules. Methods for the determination of a propensity to develop ***gastritis***, peptic ulcer disease, or ***gastric*** cancer is provided for by detection methods. Methods are also provided for the use of Hp CoA-t, thiolase or PHB synthase proteins and fragments retaining enzymatic activity in the identification of potential drug candidates for the treatment of some types of ***gastric*** disease. Pharmaceutical compositions containing Hp CoA-t protein fragments, antisense nucleic acids or other inhibitors of Hp CoA-t, thiolase and PHB synthase as well as methods for their use in the treatment of some types of ***gastric*** disease are also provided.

L11 ANSWER 139 OF 184 USPATFULL

AN 2000:31028 USPATFULL

TI ATP synthase subunit homolog

IN Tang, Y. Tom, San Jose, CA, United States Corley, Neil C., Mountain View, CA, United States Guegler, Karl J., Menlo Park, CA, United States Baughn, Mariah R., San Leandro, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 6036954 20000314

AI US 1999-373029 19990811 (9)

RLI Division of Ser. No. US 1998-154802, filed on 17 Sep 1998

DT Utility

EXNAM Primary Examiner: Achutamurthy, Ponnathapu; Assistant Examiner: Slobodyansky, Elizabeth

LREP Muenzen, Colette C.Incyte Pharmaceuticals, Inc.

CLMN Number of Claims: 2

ECL Exemplary Claim: 1

DRWN 5 Drawing Figure(s); 5 Drawing Page(s)

LN.CNT 2512

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides a human ATP synthase subunit homolog (ASYNT) and polynucleotides which identify and encode ASYNT. The invention also

provides expression vectors, host cells, ***antibodies***, agonists, and antagonists. The invention also provides methods for diagnosing, treating or preventing disorders associated with expression of ASYNT.

L11 ANSWER 140 OF 184 USPATFULL

AN 2000:1708 USPATFULL

TI Process for detecting a variant CD44 gene product

IN Ponta, Helmut, Linkenheim-Hochstetten, Germany, Federal Republic of Heider, Karl-Heinz, Waldbronn-Reichenbach, Germany, Federal Republic of Herrlich, Peter, Karlsruhe, Germany, Federal Republic of Pals, Steven T., Amsterdam, Netherlands

Dall, Peter, Dusseldorf, Germany, Federal Republic of

PA Boehringer Ingelheim International GmbH, Germany, Federal Republic of (non-U.S. corporation)

PI US 6010865 20000104

WO 9500851 19950105

AI US 1996-564225 19960603 (8)

WO 1994-EP1952 19940615

19960603 PCT 371 date

19960603 PCT 102(e) date

PRAI DE 1993-4320624 19930622

DE 1993-4320623 19930622

DE 1993-4321944 19930702

DE 1994-4414787 19940428

DT Utility

EXNAM Primary Examiner: Hutzell, Paula K.; Assistant Examiner: Ungar, Susan

LREP Finnegan, Henderson, Farabow, Garrett & Dunner, L.L.P.

CLMN Number of Claims: 7

ECL Exemplary Claim: 1,3

DRWN 45 Drawing Figure(s); 16 Drawing Page(s)

LN.CNT 1294

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention relates to a process for diagnosing and analysing tumours which is based on detecting the expression of certain variant exons of the CD44-gene. Detection may be carried out at the protein or nucleic acid level. In a preferred embodiment the expression is detected in biopsy material using exon-specific ***antibodies***. Thus, for example, v6-expression is a suitable prognostic parameter for breast cancer, the expression of a transitional epitope which is coded by exons v7 and v8 serves to diagnose cervical cancer.

L11 ANSWER 141 OF 184 USPATFULL

AN 1999:167130 USPATFULL

TI Treatment and prevention of helicobacter infection

IN Doidge, Christopher V., Vincent, Australia

Lee, Adrian, Lane Cove, Australia

Radcliff, Flona J., Sydney, Australia

Hazell, Stuart L., Glenfield, Australia

PA The University of New South Wales, Kensington, Australia (non-U.S. corporation)

CSL Limited, Victoria, Australia (non-U.S. corporation)

PI US 6005090 19991221

AI US 1996-695987 19960815 (8)

RLI Continuation-in-part of Ser. No. WO 1995-AU335, filed on 8 Jun 1995

PRAI AU 1994-6124 19940608

DT Utility

EXNAM Primary Examiner: Chin, Christopher L.; Assistant Examiner: Portner, Ginny Allen

CLMN Number of Claims: 13

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 1390

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB An antigenic preparation for use in the treatment or prevention of Helicobacter infection in a mammalian host, comprises the catalase enzyme of Helicobacter bacteria, particularly the catalase enzyme of H. ***pylori*** or H. felis, or an immunogenic fragment thereof.

L11 ANSWER 142 OF 184 USPATFULL

AN 1999:166833 USPATFULL

TI Human lysophospholipase

IN Hillman, Jennifer L., Mountain View, CA, United States
 Shah, Purvi, Sunnyvale, CA, United States
 Corley, Neil C., Mountain View, CA, United States
 Murry, Lynn E., Portola Valley, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 6004792 19991221

AI US 1998-216001 19981217 (9)

RLI Division of Ser. No. US 1997-878862, filed on 19 Jun 1997

DT Utility

EXNAM Primary Examiner: Achutamurthy, Ponnathapu; Assistant Examiner: Mayhew, Bradley S.

LREP Incyte Pharmaceuticals, Inc.

CLMN Number of Claims: 2

ECL Exemplary Claim: 1

DRWN 5 Drawing Figure(s); 5 Drawing Page(s)

LN.CNT 2205

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides a new human lysophospholipase (IHLP) and polynucleotides which identify and encode IHLP. The invention also provides expression vectors, host cells, agonists, ***antibodies*** and antagonists. The invention also provides methods for treating inflammation and disorders associated with expression of IHLP.

L11 ANSWER 143 OF 184 USPATFULL

AN 1999:166395 USPATFULL

TI IceA gene and related methods

IN Miller, Geraldine G., Franklin, TN, United States Peek, Jr., Richard M., Nashville, TN, United States Thompson, Stuart A., Whites Creek, TN, United States Blaser, Martin J., Nashville, TN, United States

PA Vanderbilt University, Nashville, TN, United States (U.S. corporation)

PI US 6004354 19991221

AI US 1998-60584 19980415 (9)

RLI Division of Ser. No. US 1996-650528, filed on 20 May 1996, now patented, Pat. No. US 5780278

DT Utility

EXNAM Primary Examiner: Chin, Christopher L.; Assistant Examiner: Portner, Ginny Allen

LREP Needle & Rosenberg
CLMN Number of Claims: 2
ECL Exemplary Claim: 1
DRWN 8 Drawing Figure(s); 7 Drawing Page(s)
LN.CNT 2180
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A purified IceA protein of Helicobacter ***pylori*** is provided. The protein is expressed as either an IceA 1 or an IceA 2 variant. A purified polypeptide fragment of the IceA protein is also provided. An antigenic fragment of IceA is provided. An isolated nucleic acid that encodes an IceA protein of H. ***pylori*** is provided. A nucleic acid that encodes an IceA 1 variant and a nucleic acid that encodes an IceA 2 variant is also provided. Fragments of the iceA gene are provided. A method of detecting the presence of an ***antibody*** against H. ***pylori*** in a sample is provided. The method comprises the following steps: a) contacting the sample with a purified IceA protein of H. ***pylori*** or a H. ***pylori*** -specific fragment thereof, and b) detecting the binding of the ***antibody*** in the sample to the protein or fragment, the detection of binding indicating the presence in the sample of ***antibodies*** against H. ***pylori*** . A method of detecting the presence of an ***antibody*** against an ulcerative Helicobacter ***pylori*** strain in a sample is also provided.

L11 ANSWER 144 OF 184 USPATFULL

AN 1999:163449 USPATFULL

TI Human myosin heavy chain-like proteins

IN Bandman, Olga, Mountain View, CA, United States Yue, Henry, Sunnyvale, CA, United States Corley, Neil C., Mountain View, CA, United States Shah, Purvi, Sunnyvale, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 6001593 19991214

AI US 1997-966318 19971107 (8)

DT Utility

EXNAM Primary Examiner: Carlson, Karen Cochrane

LREP Incyte Pharmaceuticals, Inc.

CLMN Number of Claims: 8

ECL Exemplary Claim: 1

DRWN 13 Drawing Figure(s); 13 Drawing Page(s)

LN.CNT 2545

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides human myosin heavy chain-like proteins (MHCP) and polynucleotides which identify and encode MHCP. The invention also provides expression vectors, host cells, ***antibodies***, agonists, and antagonists. The invention also provides methods for treating disorders associated with expression of MHCP.

L11 ANSWER 145 OF 184 USPATFULL

AN 1999:150966 USPATFULL

TI Human ankyrin family protein

IN Tang, Y. Tom, San Jose, CA, United States Guegler, Karl J., Menlo Park, CA, United States Corley, Neil C., Mountain View, CA, United States Yue, Henry, Sunnyvale, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 5989863 19991123

AI US 1998-172977 19981014 (9)

DT Utility

EXNAM Primary Examiner: Campbell, Eggerton A.; Assistant Examiner: Srivastava, Devesa

LREP Incyte Pharmaceuticals, Inc.

CLMN Number of Claims: 9

ECL Exemplary Claim: 1

DRWN 2 Drawing Figure(s); 5 Drawing Page(s)

LN.CNT 2663

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides a human ankyrin family protein (ANFP) and polynucleotides which identify and encode ANFP. The invention also provides expression vectors, host cells, ***antibodies***, agonists, and antagonists. The invention also provides methods for diagnosing, treating, or preventing disorders associated with expression of ANFP.

L11 ANSWER 146 OF 184 USPATFULL

AN 1999:150943 USPATFULL

TI Estimation of active infection by heliobacter ***pylori***

IN D'Angelo, Joseph P., Miami, FL, United States Zhe, Jin, Miami, FL, United States

PA Americare International Diagnostics, Inc., Miami, FL, United States (U.S. corporation)

PI US 5989840 19991123

AI US 1997-865089 19970529 (8)

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Lerner, Herbert L.; Greenberg, Laurence A.

CLMN Number of Claims: 17

ECL Exemplary Claim: 1

DRWN 9 Drawing Figure(s); 3 Drawing Page(s)

LN.CNT 586

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Disclosed is a diagnostic apparatus for estimating an active Heliobacter

pylori infectious agent in saliva, comprising in combination an

immunoassay chamber in which a first portion of said saliva is subjected to serological test for ***antibody*** to said infectious agent and a chemical reaction chamber in which a second portion of said saliva is subjected to chemical analysis for an ammonia constituent thereof.

L11 ANSWER 147 OF 184 USPATFULL

AN 1999:150925 USPATFULL

TI ATP synthase subunit homolog

IN Tang, Y. Tom, San Jose, CA, United States Corley, Neil C., Mountain View, CA, United States Guegler, Karl J., Menlo Park, CA, United States Baughn, Mariah R., San Leandro, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 5989822 19991123

AI US 1998-154802 19980917 (9)

DT Utility

EXNAM Primary Examiner: Achutamurthy, Ponnathapura; Assistant Examiner: Slobodyansky, Elizabeth

LREP Incyte Pharmaceuticals, Inc.; Muenzen, Colette C.

CLMN Number of Claims: 9

ECL Exemplary Claim: 1

DRWN 5 Drawing Figure(s); 5 Drawing Page(s)

LN.CNT 2446

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides a human ATP synthase subunit homolog (ASYNT) and polynucleotides which identify and encode ASYNT. The invention also provides expression vectors, host cells, ***antibodies***, agonists, and antagonists. The invention also provides methods for diagnosing, treating or preventing disorders associated with expression of ASYNT.

L11 ANSWER 148 OF 184 USPATFULL

AN 1999:146266 USPATFULL

TI Platinum-containing compounds, methods for their preparation and applications thereof

IN Houthoff, Hendrik J., Amsterdam, Netherlands

Reedijk, Jan, Leiden, Netherlands

Jelsma, Tinka, Almere, Netherlands

Van Es, Remco Maria, Zaan, Netherlands

van den Berg, Franciscus Michiel, Hoofddorp, Netherlands

Lempers, Edwin Leo Mario, Julianadorp, Netherlands

Bloemink, Marieke Johanna, Oegstgeest, Netherlands

PA Kreatech Diagnostics, Amsterdam, Netherlands (non-U.S. corporation)

PI US 5985566 19991116

AI US 1997-910070 19970812 (8)

RLI Continuation of Ser. No. US 1995-470265, filed on 6 Jun 1995, now patented, Pat. No. US 5714327, issued on 3 Feb 1998 which is a continuation-in-part of Ser. No. US 1993-975586, filed on 19 Oct 1993, now patented, Pat. No. US 5580990, issued on 3 Dec 1996

PRAI NL 1990-1639 19900719 .

DT Utility

EXNAM Primary Examiner: Degen, Nancy; Assistant Examiner: Sandals, William

LREP Hoffmann & Baron, LLP CLMN Number of Claims: 15 ECL Exemplary Claim: 1 DRWN No Drawings LN.CNT 1534

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides platinum-based probe compounds having the structure: ##STR1## wherein: Pt is a platinum atom, PROBE is a probe biomolecule for associating to a target biomolecule, M is a detectable marker moiety, and X and Y are stabilizing substituents. Also provided are platinum-based labeling compounds having the structure: ##STR2## wherein: Pt is a platinum atom, M is a detectable marker moiety, A is a displaceable leaving group, and X and Y are stabilizing substituents. The invention further provides platinum-based linker compounds having the structure: ##STR3## wherein: Pt is a platinum atom, A and B are the same or different reactive moieties, and X and Y are stabilizing substituents. Other Pt.sup.II and Pt.sup.IV compounds are also provided.

Moreover, the invention provides methods for the preparation and use of these compounds, as well as diagnostic kits which contain the compounds.

L11 ANSWER 149 OF 184 USPATFULL

AN 1999:136950 USPATFULL

TI Human reticulocalbin isoforms

IN Bandman, Olga, Mountain View, CA, United States Hillman, Jennifer L., Mountain View, CA, United States Lal, Preeti, Santa Clara, CA, United States Corley, Neil C., Mountain View, CA, United States Shah, Purvi, Sunnyvale, CA, United States

PA Incyte Pharamceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 5976801 19991102

AI US 1997-910927 19970808 (8)

DT Utility

EXNAM Primary Examiner: Fredman, Jeffrey

LREP Price, Leanne C. Incyte Pharmaceuticals, Inc.

CLMN Number of Claims: 18

ECL Exemplary Claim: 1

DRWN 18 Drawing Figure(s); 18 Drawing Page(s)

LN.CNT 2648

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides two human reticulocalbin isoforms designated individually as RCN .gamma. and RCN .delta. and collectively as RCN, and polynucleotides which identify and encode RCN. The invention also provides expression vectors, host cells, agonists, ***antibodies*** and antagonists. The invention also provides methods for treating disorders associated with expression of RCN.

L11 ANSWER 150 OF 184 USPATFULL

AN 1999:124759 USPATFULL

TI Human lysophospholipase

IN Hillman, Jennifer L., Mountain View, CA, United States Shah, Purvi, Sunnyvale, CA, United States Murry, Lynn E., Portola Valley, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 5965423 19991012

AI US 1998-22940 19980212 (9)

RLI Continuation-in-part of Ser. No. US 1997-844120, filed on 29 Apr 1997

DT Utility

EXNAM Primary Examiner: Hendricks, Keith D.; Assistant Examiner: Mayhew, Bradley S.

LREP Mohan-Peterson, SheelaIncyte Pharmaceuticals, Inc.

CLMN Number of Claims: 9

ECL Exemplary Claim: 1

DRWN 9 Drawing Figure(s); 9 Drawing Page(s)

LN.CNT 2264

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides a human lysophospholipase (NHLP) and polynucleotides which identify and encode NHLP. The invention also provides expression vectors, host cells, ***antibodies***, agonists, and antagonists. The invention also provides methods for treating or preventing disorders associated with expression of NHLP.

L11 ANSWER 151 OF 184 USPATFULL

AN 1999:121547 USPATFULL

TI ATP synthase Fo subunit

IN Hillman, Jennifer L., Mountain View, CA, United States Goli, Surya K., Sunnyvale, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 5962646 19991005

AI US 1998-216625 19981216 (9)

RLI Division of Ser. No. US 1998-94080, filed on 9 Jun 1998 which is a division of Ser. No. US 1997-948195, filed on 9 Oct 1997, now patented, Pat. No. US 5763248 which is a continuation of Ser. No. US 1997-819395, filed on 17 Mar 1997, now abandoned

DT Utility

EXNAM Primary Examiner: Degen, Nancy; Assistant Examiner: Wang, Andrew LREP Mohan-Peterson, SheelaIncyte Pharmaceuticals, Inc.

CLMN Number of Claims: 2

ECL Exemplary Claim: 1

DRWN 4 Drawing Figure(s); 4 Drawing Page(s)

LN.CNT 1999

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides a human ATP synthase d subunit (ASYSD) and polynucleotides which encode ASYSD. The invention also provides expression vectors, host cells, agonists, antisense molecules, ***antibodies***, or antagonists. The invention also provides to methods for producing ASYSD and for treating disorders associated with expression of ASYSD.

L11 ANSWER 152 OF 184 USPATFULL

AN 1999:106303 USPATFULL

TI Method of screening for ATP synthase Fo subunit

IN Hillman, Jennifer L., Mountain View, CA, United States Goli, Surya K., Sunnyvale, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 5948625 19990907

AI US 1998-94080 19980609 (9)

RLI Division of Ser. No. US 1997-948195, filed on 9 Oct 1997, now patented, Pat. No. US 5763248 which is a continuation of Ser. No. US 1997-819395, filed on 17 Mar 1997, now abandoned

DT Utility

EXNAM Primary Examiner: LeGuyader, John L., Assistant Examiner: Wang, Andrew

LREP Mohan-Peterson, SheelaIncyte Pharmaceuticals, Inc.

CLMN Number of Claims: 4

ECL Exemplary Claim: 1

DRWN 6 Drawing Figure(s); 6 Drawing Page(s)

LN.CNT 1993

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides a human ATP synthase d subunit (ASYSD) and polynucleotides which encode ASYSD. The invention also provides expression vectors, host cells, agonists, antisense molecules, ***antibodies***, or antagonists. The invention also provides methods for producing ASYSD and for treating disorders associated with expression of ASYSD.

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L11 ANSWER 153 OF 184 USPATFULL
AN 1999:89021 USPATFULL
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TI Vacuolar ***atpase*** subunit AC45

IN Hillman, Jennifer L., Mountain View, CA, United States Shah, Purvi, Sunnyvale, CA, United States Corley, Neil C., Mountain View, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 5932444 19990803

AI US 1997-959011 19971028 (8)

DT Utility

EXNAM Primary Examiner: Huff, Sheela; Assistant Examiner: Eyler, Yvonne

LREP Incyte Pharmaceuticals, Inc.

CLMN Number of Claims: 8

ECL Exemplary Claim: 1

DRWN 10 Drawing Figure(s), 10 Drawing Page(s)

LN.CNT 2221

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides a human vacuolar ***ATPase*** subunit AC45 (HAC45) and polynucleotides which identify and encode HAC45. The invention also provides expression vectors, host cells, agonists, ***antibodies***, and antagonists. The invention also provides methods for treating or preventing disorders associated with expression of HAC45.

L11 ANSWER 154 OF 184 USPATFULL

AN 1999:89007 USPATFULL

TI ***Immunoassay*** for H. ***pylori*** in fecal specimens

IN Larka, Christopher Vance, Cincinnati, OH, United States
Yi, Ching Sui Arthur, Cincinnati, OH, United States
Kozak, Kenneth James, Cincinnati, OH, United States

PA Meridian Diagnostics, Inc., Cincinnati, OH, United States (U.S. corporation)

PI US 5932430 19990803

AI US 1998-37894 19980310 (9)

RLI Continuation-in-part of Ser. No. US 1997-897732, filed on 21 Jul 1997, now patented, Pat. No. US 5871942 which is a continuation-in-part of Ser. No. US 1996-647115, filed on 9 May 1996, now patented, Pat. No. US 5716791

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Thompson, Hine & Flory LLP

CLMN Number of Claims: 20

ECL Exemplary Claim: 1

DRWN 2 Drawing Figure(s); 1 Drawing Page(s)

LN.CNT 616

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A process for the determination of H. ***pylori*** in a fecal specimen comprising (a) dispersing a fecal specimen suspected of carrying H. ***pylori*** in a sample diluent; (b) contacting the fecal specimen in the diluent with a first polyclonal ***antibody*** for H. ***pylori*** antigen to form a complex of the ***antibody*** and the antigen; (c) separating said specimen and said

complex; (d) exposing the complex to a second polyclonal
antibody for said antigen and a portion of the ***antibody***
reacting with said complex, one of said first and second
antibody being bound to a solid carrier and the other being
labeled with a detection agent; and (e) determining the amount of the
labeled ***antibody*** and in turn determining the presence of H.
pylori antigen in said fecal specimen.

L11 ANSWER 155 OF 184 USPATFULL

AN 1999:85216 USPATFULL

TI Compositions comprising isolated Helicobacter ***pylori*** CagI polynucleotides and method of preparation thereof

IN Covacci, Antonello, Siena, Italy

PA Chiron S.p.A., Italy (non-U.S. corporation)

PI US 5928865 19990727

AI US 1995-477451 19950607 (8)

RLI Continuation-in-part of Ser. No. US 1995-425194, filed on 20 Apr 1995, now abandoned And Ser. No. US 1995-471491, filed on 6 Jun 1995 which is a division of Ser. No. US 256848

PRAI IT 1992-FI52 19920302

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Woodcock, Washburn, Kurtz, Mackiewicz & Norris, Harbin, Alisa A.; Blackburn, Robert P.

CLMN Number of Claims: 6

ECL Exemplary Claim: 1

DRWN 4 Drawing Figure(s), 120 Drawing Page(s)

LN.CNT 6155

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Helicobacter ***pylori*** is known to cause or be a cofactor in type B ***gastritis*** , peptic ulcers, and ***gastric*** tumors. In both developed and developing countries, a high percentage of people are infected with this bacterium. The present invention relates generally to a certain H. ***pylori*** region located 5' to the CagA gene locus, to proteins encoded thereby, and to the use of these genes and proteins for diagnostic and vaccine applications.

L11 ANSWER 156 OF 184 USPATFULL

AN 1999:43190 USPATFULL

TI Method for stimulating production of variable region gene family restricted ***antibodies*** through B-cell superantigen vaccination

IN Silverman, Gregg J., Encinitas, CA, United States

PA The Regents of the University of California, Oakland, CA, United States (U.S. corporation)

PI US 5891438 19990406

WO 9409818 19940511

AI US 1995-428197 19950714 (8)

WO 1993-US10555 19931029

19950714 PCT 371 date

19950714 PCT 102(e) date

RLI Continuation-in-part of Ser. No. US 1992-969936, filed on 30 Oct 1992, now abandoned

DT Utility

EXNAM Primary Examiner: Feisee, Lila; Assistant Examiner: Ungar, Susan

LREP Fish & Richardson P.C.
CLMN Number of Claims: 3
ECL Exemplary Claim: 1,3
DRWN 29 Drawing Figure(s); 12 Drawing Page(s)
LN.CNT 2731
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Criteria for identifying potential B cell superantigens are disclosed, together with a method for determining whether these candidate antigens have B cell superantigenic activity. Methods for constructing and using a vaccine including B cell superantigens are also disclosed. Identification is based on characterizing the structure of Ig binding sites which interact with the candidate antigen assessment of Ig V region diversity on binding of candidate and conventional antigens, confirmation of sAg activity in interactions between candidate antigens and whole cells, confirmation of whether the candidate antigen induces B cell mitogenesis, determination of the earliest point in B cell development where cellular co-factors are required for sAg activity and, for reference, determination of V region usage in responder populations. Once a B cell superantigen is characterized, it is purified and conjugated by chemical means to a polysaccharide or glycoprotein component from a microbial capsule, cell wall, envelope or other component preferably using components which stimulate production of ***antibodies*** with the same V region restriction as ***antibodies*** whose production is stimulated by the B cell superantigen.

L11 ANSWER 157 OF 184 USPATFULL

AN 1999:40212 USPATFULL

TI ATP-dependent RNA helicase protein

IN Bandman, Olga, Mountain View, CA, United States Guegler, Karl J., Menlo Park, CA, United States Corley, Neil C., Mountain View, CA, United States Shah, Purvi, Sunnyvale, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 5888792 19990330

AI US 1997-892256 19970711 (8)

DT Utility

EXNAM Primary Examiner: Wax, Robert A.; Assistant Examiner: Slobodyomsky, Elijobette

LREP Incyte Pharmaceuticals, Inc.

CLMN Number of Claims: 9

ECL Exemplary Claim: 1

DRWN 14 Drawing Figure(s); 14 Drawing Page(s)

LN.CNT 2270

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides a human ATP-dependent RNA helicase (ADRH-1) and polynucleotides which identify and encode ADRH-1. The invention also provides expression vectors, host cells, agonists, ***antibodies*** and antagonists. The invention also provides methods for treating disorders associated with expression of ADRH-1.

L11 ANSWER 158 OF 184 USPATFULL

AN 1999:30582 USPATFULL

TI Methods and compositions for the diagnosis of extraesophageal

gastric reflux

IN Koufman, James, Winston-Salem, NC, United States

PA Wake Forest University, Winston-Salem, NC, United States (U.S. corporation)

PI US 5879897 19990309

AI US 1996-717793 19960923 (8)

DT Utility

EXNAM Primary Examiner: Saunders, David

LREP Arnold, White & Durkee CLMN Number of Claims: 23 ECL Exemplary Claim: 1

DRWN 3 Drawing Figure(s); 2 Drawing Page(s)

LN.CNT 1865

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Disclosed are methods of detecting a ***gastric*** reflux in the esophagus or in the throat of a subject. The basis of the method is the detection of the presence of pepsin or ***pepsinogen*** at higher than normal levels. Detection is preferably by an ***immunoassay*** technique.

L11 ANSWER 159 OF 184 USPATFULL

AN 1999:27407 USPATFULL

TI Helicobacter TagA gene fusion protein

IN Cover, Timothy L., Nashville, TN, United States
 Blaser, Martin J., Nashville, TN, United States
 Kleanthous, Harry, Cambridge, MA, United States
 Tummuru, Murali K. R., Nashville, TN, United States

PA Vanderbilt University, Nashville, TN, United States (U.S. corporation)

PI US 5876943 19990302

AI US 1998-34306 19980302

RLI Continuation of Ser. No. US 1994-316397, filed on 30 Sep 1994, now patented, Pat. No. US 5733340 which is a continuation-in-part of Ser. No. US 1993-53614, filed on 26 Apr 1993, now patented, Pat. No. US 5403924, issued on 4 Apr 1995 which is a continuation-in-part of Ser. No. US 1992-959940, filed on 13 Oct 1992, now abandoned

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Needle & Rosenberg

CLMN Number of Claims: 2

ECL Exemplary Claim: 1

DRWN 4 Drawing Figure(s); 4 Drawing Page(s)

LN.CNT 2470

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides an isolated nucleic acid encoding an approximately 120-128 kilodalton antigen of Helicobacter ***pylori***, or an antigenic fragment thereof, wherein the antigen is associated with peptic ulceration. The present invention also provides methods of detecting the presence of a Helicobacter ***pylori*** strain possessing the 120-128 kilodalton antigen in a subject, comprising the steps of contacting an ***antibody*** -containing sample from the subject with a detectable amount of the tagA antigen or antigenic polypeptide of the present invention and detecting the binding of the antigen or fragment and the ***antibody***. The detection of a strain expressing the TagA antigen is an indication of predisposition to

peptic ulceration and ****gastric*** carcinoma. A mutant H. ***pylori*** not expressing a functional TagA antigen is also provided.

L11 ANSWER 160 OF 184 USPATFULL

AN 1999:21925 USPATFULL

II ***Immunoassay*** for H. ***pylori*** in fecal specimens

IN Larka, Christopher Vance, Cincinnati, OH, United States Yi, Ching Sui Arthur, Cincinnati, OH, United States Kozak, Kenneth James, Cincinnati, OH, United States

PA Meridian Diagnostics, Inc., Cincinnati, OH, United States (U.S. corporation)

PI US 5871942 19990216

AI US 1997-897732 19970721 (8)

RLI Continuation-in-part of Ser. No. US 1996-647115, filed on 9 May 1996, now patented, Pat. No. US 5716791

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Thompson Hine & Flory LLP

CLMN Number of Claims: 14

ECL Exemplary Claim: 1

DRWN 2 Drawing Figure(s), 1 Drawing Page(s)

LN.CNT 502

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A process for the determination of H. ***pylori*** in a fecal specimen which comprises; (a) collecting a smear of a fecal specimen on a substrate; (b) immersing at least the portion of the substrate carrying the smear in a sample diluent so as to disperse the fecal specimen in the diluent; (c) contacting the fecal specimen in the diluent with a first polyclonal ***antibody*** for H. ***pylori*** antigen to form a complex of the ***antibody*** and the antigen; (d) separating the specimen and the complex; (e) exposing the complex to a second polyclonal ***antibody*** for the antigen and a portion of the ***antibody*** reacting with the complex, one of said first and second ***antibody*** being bound to a solid carrier and the other being labelled with a detection agent; and (f) determining the amount of the labelled ***antibody*** and in turn determining the presence of H. ***pylori*** antigen in the fecal specimen.

L11 ANSWER 161 OF 184 USPATFULL

AN 1999:4417 USPATFULL

TI Human lysophospholipase

IN Hillman, Jennifer L., Mountain View, CA, United States Shah, Purvi, Sunnyvale, CA, United States Murry, Lynn E., Portola Valley, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 5858756 19990112

AI US 1997-844120 19970429 (8)

DT Utility

EXNAM Primary Examiner: Hendricks, Keith D.; Assistant Examiner: Mayhew, Bradley S.

LREP Incyte Pharmaceuticals, Inc.; Billings, Lucy J.; Mohan-Peterson, Sheela CLMN Number of Claims: 8

ECL Exemplary Claim: 1

DRWN 8 Drawing Figure(s); 8 Drawing Page(s)

LN.CNT 1990

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides a human lysophospholipase (NHLP) and polynucleotides which identify and encode NHLP. The invention also provides expression vectors, host cells, agonists, ***antibodies*** and antagonists. In addition, the invention provides methods for producing NHLP and for treating or preventing disorders associated with expression of NHLP.

L11 ANSWER 162 OF 184 USPATFULL

AN 1998:154068 USPATFULL

TI Test kits and methods for detecting H. ***pylori***

IN Pronovost, Allan David, San Diego, CA, United States Pawlak, Jan Waclaw, Cardiff, CA, United States Condon, Kristy S., San Diego, CA, United States

PA Quidel Corporation, San Diego, CA, United States (U.S. corporation)

PI US 5846751 19981208

AI US 1995-486843 19950607 (8)

RLI Division of Ser. No. US 1994-292932, filed on 18 Aug 1994 which is a continuation of Ser. No. US 1993-22817, filed on 24 Feb 1993, now abandoned which is a continuation of Ser. No. US 1990-621845, filed on 4 Dec 1990, now abandoned

DT Utility

EXNAM Primary Examiner: Loring, Susan A.

CLMN Number of Claims: 6

ECL Exemplary Claim: 1

DRWN 1 Drawing Figure(s); 1 Drawing Page(s)

LN.CNT 725

AB A sensitive and specific antigen-preparation for the detection of Helicobacter ***pylori*** in biological samples is disclosed. The preparation uses a range of antigens derived from size exclusion chromatography of detergent-solubilized H. ***pylori*** cells. Serological assays such as ELISA, latex agglutination, and rapid EIA assays utilizing the improved antigen preparation, and a kit for use in these serological assays are also disclosed.

L11 ANSWER 163 OF 184 USPATFULL

AN 1998:143906 USPATFULL

TI Helicobacter ***pylori*** haemagglutinin protease protein, nucleic acid encoding therefor and ***antibodies*** specific thereto

IN Smith, Andrew William, Kent, Great Britain

PA Reckitt & Colman Products Limited, London, United Kingdom (non-U.S. corporation)

PI US 5837502 19981117

WO 9501445 19950112

AI US 1996-578516 19960916 (8)

WO 1994-GB1406 19940629

19960916 PCT 371 date

19960916 PCT 102(e) date

PRAI GB 1993-13437 19930630

DT Utility

EXNAM Primary Examiner: Horlick, Kenneth R.; Assistant Examiner: Tung, Joyce LREP Wolf, Greenfield & Sacks, P.C.

CLMN Number of Claims: 18

ECL Exemplary Claim: 1

DRWN 2 Drawing Figure(s); 2 Drawing Page(s)

LN.CNT 714

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Helicobacter ***pylori*** (H. ***pylori***)

haemagglutinin/protease protein, nucleic acids encoding therefor and ****antibodies*** specific thereto are described and, in particular, to their use in the identification of H. ***pylori*** and in the diagnosis of H. ***pylori*** infection. Also described are kits for the identification and diagnosis of H. ***pylori*** infection.

L11 ANSWER 164 OF 184 USPATFULL

AN 1998:122236 USPATFULL

TI DNA encoding a histamine H2 receptor

IN Murry, Lynn E., Portola Valley, CA, United States Au-Young, Janice, Berkeley, CA, United States Guegler, Karl J., Menlo Park, CA, United States Goli, Surya K., Sunnyvale, CA, United States

PA Incyte Pharmaceuticals, Palo Alto, CA, United States (U.S. corporation)

PI US 5817480 19981006

AI US 1996-748485 19961107 (8)

DT Utility

EXNAM Primary Examiner: Walsh, Stephen; Assistant Examiner: Basham, Daryl A.

LREP Billings, Lucy J.

CLMN Number of Claims: 7

ECL Exemplary Claim: 1

DRWN 14 Drawing Figure(s); 14 Drawing Page(s)

LN.CNT 2287

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides a novel histamine H2 receptor (H2RH) and polynucleotides which identify and encode H2RH. The invention also provides genetically engineered expression vectors and host cells comprising the nucleic acid sequences encoding H2RH and a method for producing H2RH. The invention also provides for agonists, ***antibodies***, or antagonists specifically binding H2RH, and their use, in the prevention and treatment of diseases in which H2RH is expressed. Additionally, the invention provides for the use of antisense molecules to polynucleotides encoding H2RH for the treatment of diseases associated with the expression of H2RH. The invention also provides diagnostic assays which utilize the polynucleotide, or fragments or the complement thereof, and ***antibodies*** specifically binding H2RH.

L11 ANSWER 165 OF 184 USPATFULL

AN 1998:118977 USPATFULL

TI Antigen preparation for detecting H. ***pylori***

IN Pronovost, Allan David, San Diego, CA, United States Pawlak, Jan Waclaw, Cardiff, CA, United States Condon, Kristy S., San Diego, CA, United States

PA Quidel Corporation, San Diego, CA, United States (U.S. corporation)

PI US 5814455 19980929

AI US 2929325 19940818 (8)

RLI Continuation of Ser. No. 22817, filed on 24 Feb 1993, now abandoned which is a continuation of Ser. No. 621845, filed on 4 Dec 1990, now abandoned

DT Utility

EXNAM Primary Examiner: Sidberry, Hazel F.

LREP Morrison & Foerster CLMN Number of Claims: 2

ECL Exemplary Claim: 1

DRWN 1 Drawing Figure(s); 1 Drawing Page(s)

LN.CNT 730

AB A sensitive and specific antigen preparation for the detection of Helicobacter ***pylori*** in biological samples is disclosed. The preparation uses a range of antigens derived from size exclusion chromatography of detergent-solubilized H. ***pylori*** cells. Serological assays such as ELISA, latex agglutination, and rapid EIA assays utilizing the improved antigen preparation, and a kit for use in these serological assays are also disclosed.

L11 ANSWER 166 OF 184 USPATFULL

AN 1998:111817 USPATFULL

TI GDP-L-fucose: .beta.-D-galactoside 2-.alpha.-L-fucosyltransferases, DNA sequences encoding the same, method for producing the same and a method of genotyping a person

IN Lowe, John B., 3125 Bolgos Cir., Ann Arbor, MI, United States 48105 Lennon, Gregory, 8309 Norris Canyon, Castro Valley, CA, United States 94552

Rouquier, Sylvie, 5, rue du Cannau, 34000 Montpellier, France Giorgi, Dominique, 5, rue du Cannau, 34000 Montpellier, France Kelly, Robert J., 3164 Concord, Trenton, MI, United States 48183

PI US 5807732 19980915

AI US 1995-395800 19950228 (8)

DT Utility

EXNAM Primary Examiner: Wax, Robert A., Assistant Examiner: Nashed, Nashaat T.

LREP Oblon, Spivak, McClelland, Maier & Neustadt, P.C.

CLMN Number of Claims: 12

ECL Exemplary Claim: 9

DRWN 30 Drawing Figure(s); 23 Drawing Page(s)

LN.CNT 2647

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The gene encoding GDP-L-fucose: .beta.-D-Galactoside
2-.alpha.-L-fucosyltransferase has been cloned, and a mutation in this gene has been found to be responsible for an individual being a non-secretor.

L11 ANSWER 167 OF 184 USPATFULL

AN 1998:88644 USPATFULL

TI F.sub.0 ATP synthase subunit

IN Hillman, Jennifer L., Mountain View, CA, United States Goli, Surya K., Sunnyvale, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 5786150 19980728

AI US 1997-815177 19970311 (8)

DT Utility

EXNAM Primary Examiner: Patterson, Jr., Charles L.

LREP Billings, Lucy J.; Mohan-Peterson, SheelaIncyte Pharmceuticals

CLMN Number of Claims: 10 ECL Exemplary Claim: 1

DRWN 4 Drawing Figure(s); 3 Drawing Page(s)

LN.CNT 1940

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides a human ATP synthase subunit (ASYS) and polynucleotides which encode ASYS. The invention also provides expression vectors, host cells, agonists, antisense molecules, ***antibodies***, or antagonists. The invention also provides methods for producing ASYS and for treating disorders associated with expression of ASYS.

L11 ANSWER 168 OF 184 USPATFULL

AN 1998:82577 USPATFULL

TI IceA gene and related methods

IN Miller, Geraldine G., Franklin, TN, United States Peek, Jr., Richard M., Nashville, TN, United States Thompson, Stuart A., Whites Creek, TN, United States Blaser, Martin J., Nashville, TN, United States

PA Vanderbilt University, Nashville, TN, United States (U.S. corporation)

PI US 5780278 19980714

AI US 1996-650528 19960520 (8)

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Needle & Rosenberg, P.C.

CLMN Number of Claims: 17

ECL Exemplary Claim: 1

DRWN 8 Drawing Figure(s); 7 Drawing Page(s)

LN.CNT 2020

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A purified IceA protein of Helicobacter ***pylori*** is provided. The protein is expressed as either an IceA 1 or an IceA 2 variant. A purified polypeptide fragment of the IceA protein is also provided. An antigenic fragment of IceA is provided. An isolated nucleic acid that encodes an IceA protein of H. ***pylori*** is provided. A nucleic acid that encodes an IceA 1 variant and a nucleic acid that encodes an IceA 2 variant is also provided. Fragments of the iceA gene are provided. A method of detecting the presence of an ***antibody*** against H. ***pylori*** in a sample is provided. The method comprises the following steps: a) contacting the sample with a purified IceA protein of H. ***pylori*** or a H. ***pylori*** -specific fragment thereof, and b) detecting the binding of the ***antibody*** in the sample to the protein or fragment, the detection of binding indicating the presence in the sample of ***antibodies*** against H. ***pylori*** . A method of detecting the presence of an ***antibody*** against an ulcerative Helicobacter ***pylori*** strain in a sample is also provided.

L11 ANSWER 169 OF 184 USPATFULL

AN 1998:72414 USPATFULL

TI Methods for the diagnosis of diabetes and prediabetic conditions

IN MacKay, Ian Reay, Malvern, Australia Rowley, Merrill Joy, Camberwell, Australia Zimmet, Paul Zev, Toorak, Australia

PA Monash University, Clayton, Australia (non-U.S. corporation)

PI US 5770381 19980623

WO 9418568 19940818

AI US 1995-495584 19951010 (8)

WO 1994-AU56 19940209

19951010 PCT 371 date

19951010 PCT 102(e) date

PRAI AU 1993-7168 199

19930209

DT Utility

EXNAM Primary Examiner: Hendricks, Keith D.

LREP Foley & Lardner

CLMN Number of Claims: 21

ECL Exemplary Claim: 1

DRWN 6 Drawing Figure(s); 3 Drawing Page(s)

LN.CNT 1065

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A method for detecting autoantibodies to glutamic acid decarboxylase (GAD) in the serum of a patient as diagnostic of a diabetic or prediabetic condition in the patient, comprises contacting a serum sample from the patient with a GAD antigen and detecting binding of autoantibodies to GAD in the sample by the GAD antigen, wherein the GAD antigen comprises a GAD preparation containing an enhanced amount of dimer(s) or oligomer(s) of the 65 kD or 67 kD isoforms, or both, of GAD. A diagnostic kit is also inclosed.

L11 ANSWER 170 OF 184 USPATFULL

AN 1998:65037 USPATFULL

TI CDNA encoding a human ATP synthase Fo subunit (ASYSD)

IN Hillman, Jennifer L., Mountain View, CA, United States Goli, Surya K., Sunnyvale, CA, United States

PA Incyte Pharmaceuticals, Inc., Palo Alto, CA, United States (U.S. corporation)

PI US 5763248 19980609

AI US 1997-948195 19971009 (8)

RLI Continuation of Ser. No. US 1997-819395, filed on 17 Mar 1997, now abandoned

DT Utility

EXNAM Primary Examiner: LeGuyader, John L.; Assistant Examiner: Wang, Andrew

LREP Billings, Lucy J.

CLMN Number of Claims: 6

ECL Exemplary Claim: 1

DRWN 5 Drawing Figure(s); 4 Drawing Page(s)

LN.CNT 1963

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides a human ATP synthase d subunit (ASYSD) and polynucleotides which encode ASYSD. The invention also provides expression vectors, host cells, agonists, antisense molecules, ****antibodies****, or antagonists. The invention also provides methods for producing ASYSD and for treating disorders associated with expression of ASYSD.

L11 ANSWER 171 OF 184 USPATFULL

AN 1998:33767 USPATFULL

TI Taga gene and methods for detecting predisposition to peptic ulceration and ***gastric*** carcinoma

IN Cover, Timothy L., Nashville, TN, United States Blaser, Martin J., Nashville, TN, United States

Kleanthous, Harry, Cambridge, MA, United States Tummuru, Murali K. R., Nashville, TN, United States

PA Vanderbilt University, Nashville, TN, United States (U.S. corporation) OraVax, Inc., Cambridge, MA, United States (U.S. corporation)

PI US 5733740 19980331

AI US 1994-316397 19940930 (8)

RLI Continuation-in-part of Ser. No. US 1993-53614, filed on 26 Apr 1993, now patented, Pat. No. US 5403924 which is a continuation-in-part of Ser. No. US 1992-959940, filed on 13 Oct 1992, now abandoned

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny

LREP Needle & Rosenberg, P.C.

CLMN Number of Claims: 6

ECL Exemplary Claim: 1

DRWN 4 Drawing Figure(s); 4 Drawing Page(s)

LN.CNT 2520

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB An isolated nucleic acid encoding an approximately 120-128 kilodalton antigen of Helicobacter ***pylori***, or an antigenic fragment thereof, wherein the antigen is associated with peptic ulceration. The present invention also provides methods of detecting the presence of a Helicobacter ***pylori*** strain possessing the 120-128 kilodalton antigen in a subject, comprising the steps of contacting an ***antibody*** -containing sample from the subject with a detectable amount of the tagA antigen or antigenic polypeptide of the present invention and detecting the binding of the antigen or fragment and the ***antibody*** The detection of a strain expressing the TagA antigen is an indication of predisposition to peptic ulceration and ***gastric*** carcinoma. A mutant H. ***pylori*** not expressing a functional TagA antigen is also provided.

L11 ANSWER 172 OF 184 USPATFULL

AN 1998:14644 USPATFULL

TI ***Immunoassay*** for H. ***pylori*** in fecal specimens

IN Larka, Christopher Vance, Cincinnati, OH, United States Yi, Ching Sui Arthur, Cincinnati, OH, United States Kozak, Kenneth James, Cincinnati, OH, United States

PA Meridian Diagnostics, Inc., Cincinnati, OH, United States (U.S. corporation)

PI US 5716791 19980210

AI US 1996-647115 19960509 (8)

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Thompson Hine & Flory LLP

CLMN Number of Claims: 13

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 457

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A process for the determination of H. ***pylori*** in a fecal specimen comprising (a) dispersing a fecal specimen suspected of carrying H. ***pylori*** in a sample diluent; (b) contacting the fecal specimen in the diluent with a first polyclonal ***antibody***

for H. ***pylori*** antigen to form a complex of the

antibody and the antigen; (c) separating said specimen and said
complex; (d) exposing the complex to a second polyclonal

antibody for said antigen and a portion of the ***antibody***
reacting with said complex, one of said first and second

antibody being bound to a solid carrier and the other being
labeled with a detection agent; and (e) determining the amount of the
labeled ***antibody*** and in turn determining the presence of H.

pylori antigen in said fecal specimen.

L11 ANSWER 173 OF 184 USPATFULL

AN 1998:11876 USPATFULL

TI Platinum-containing compounds, methods for their preparation and applications thereof

IN Houthoff, Hendrik J., Amsterdam, Netherlands
 Reedijk, Jan, Leiden, Netherlands
 Jelsma, Tinka, Almere, Netherlands
 Van Es, Remco Maria, Koog a/d Zaan, Netherlands
 van den Berg, Franciscus Michiel, Hoofddorp, Netherlands
 Lempers, Edwin Leo Marlo, Julianadorp, Netherlands
 Bloemink, Marieke Johanna, Oegstgeest, Netherlands

PA Kreatech Diagnostics, Amsterdam, Netherlands (non-U.S. corporation)

PI US 5714327 19980203

AI US 1995-470265 19950606 (8)

RLI Continuation-in-part of Ser. No. US 1993-975586, filed on 29 Oct 1993, now patented, Pat. No. US 5580990, issued on 3 Dec 1996

PRAI NL 1990-1639 19900719

DT Utility

EXNAM Primary Examiner: Guzo, David; Assistant Examiner: Sandals, William

LREP Hoffmann & Baron, LLP CLMN Number of Claims: 21 ECL Exemplary Claim: 1 DRWN No Drawings

LN.CNT 1543

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The invention provides platinum-based probe compounds having the structure: ##STR1## wherein: Pt is a platinum atom, PROBE is a probe biomolecule for associating to a target biomolecule, M is a detectable marker moiety, and X and Y are stabilizing substituents. Also provided are platinum-based labeling compounds having the structure: ##STR2## wherein: Pt is a platinum atom, M is a detectable marker moiety, A is a displaceable leaving group, and X and Y are stabilizing substituents. The invention further provides platinum-based linker compounds having the structure: ##STR3## wherein: Pt is a platinum atom, A and B are the same or different reactive moieties, and X and Y are stabilizing substituents. Other Pt.sup.II and Pt.sup.IV compounds are also provided. Moreover, the invention provides methods for the preparation and use of these compounds, as well as diagnostic kits which contain the compounds.

L11 ANSWER 174 OF 184 USPATFULL

AN 97:20430 USPATFULL

TI Isolated Helicobacter hepaticus

IN Ward, Jerrold M., Gaithersburg, MD, United States Fox, James G., Harvard, MA, United States Collins, Jr., Michael J., Laurel, MD, United States Gorelick, Peter L., Frederick, MD, United States

Benveniste, Raoul E., Bethesda, MD, United States

Tully, Joseph G., Germantown, MD, United States

Gonda, Matthew A., Walkersville, MD, United States

Paster, Bruce J., Lee, NH, United States

Dewhirst, III, Floyd E., Medfield, MA, United States

PA The United States of America as represented by the Department of Health and Human Services, Washington, DC, United States (U.S. government)

PI US 5610060 19970311

AI US 1994-266414 19940624 (8)

DT Utility

EXNAM Primary Examiner: Rollins, John W.; Assistant Examiner: Ware, Deborah

LREP Needle & Rosenberg, P.C.

CLMN Number of Claims: 1

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 1816

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB An isolated bacterium of the genus Helicobacter, characterized by the 16S ribosomal RNA encoding nucleotide sequence defined in the Sequence Listing as SEQ ID NO:1 is provided. An isolated nucleic acid having the nucleotide sequence defined in the Sequence Listing as SEQ ID NO:1 is provided. Such a nucleic acid can be used for diagnosis of infection with H. hepaticus. A nucleic acid of the present invention in a vector suitable for expression of the nucleic acid is also provided. The vector can be in a host suitable for expressing the nucleic acid. A purified antigen specific for H. hepaticus is provided. A method of making an animal model for chronic Helicobacter infection is also provided.

L11 ANSWER 175 OF 184 USPATFULL

AN 96:96934 USPATFULL

TI Methods and compositions for the detection and treatment of diseases associated with antigens of microorganisms

IN Calenoff, Emanuel, Chicago, IL, United States

PA Enteron, L.P., Oak Brook, IL, United States (U.S. corporation)

PI US 5567594 19961022

AI US 1993-170017 19931220 (8)

RLI Continuation-in-part of Ser. No. US 1991-693232, filed on 26 Apr 1991, now abandoned

DT Utility

EXNAM Primary Examiner: Knode, Marian C.; Assistant Examiner: Duffy, Patricia

A.

LREP Brinks Hofer Gilson & Lione

CLMN Number of Claims: 15

ECL Exemplary Claim: 1

DRWN 5 Drawing Figure(s); 5 Drawing Page(s)

LN.CNT 1943

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A library of isolated and purified antigens specific for a microorganism is a set of individual molecules. The library forms antigen-

****antibody*** complexes useful in the context of diagnosing and treating conditions associated with a specific microorganism such as H.

pylori -induced gastro-duodenal disease. For the antigen-

antibody complexes in question the ***antibody*** is an immunoglobulin, which is IgE if the antigens are allergens. Complexes

with IgA, IgG and IgM are also useful. By this multivariate approach, a specific condition is diagnosed with high sensitivity and specificity by determining whether complexes form between a specific antigen library and a biological sample which contains immunoglobulins from an individual. Such libraries also are useful for immunotherapy.

L11 ANSWER 176 OF 184 USPATFULL

AN 96:91959 USPATFULL

TI TNF receptor-associated intracellular signaling proteins and methods of use

IN Goeddel, David V., South San Francisco, CA, United States Hsu, Hailing, South San Francisco, CA, United States

PA Tularik, Inc., So. San Francisco, CA, United States (U.S. corporation)

PI US 5563039 19961008

AI US 1995-414625 19950331 (8)

DT Utility

EXNAM Primary Examiner: Ulm, John

LREP Flehr, Hohbach, Test, Albritton & Herbert

CLMN Number of Claims: 10 ECL Exemplary Claim: 1 DRWN No Drawings

LN.CNT 1317

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A novel family of intracellular signaling proteins, exemplified by a Tumor Necrosis Factor Receptor-1 Associated Death Domain protein (TRADD), share a common TRADD sequence and include transducers of signals that modulate cell growth, differentiation and apoptosis. As such, the TRADD proteins, TRADD-encoding nucleic acids, and natural TRADD intracellular binding targets provide both important targets and means for therapeutic intervention. In particular, the invention provides isolated TRADDs and TRADD fragments, nucleic acids encoding the subject TRADDs and TRADD fragments or capable of selectively hybridizing to such TRADD-encoding nucleic acids, vectors and cells comprising TRADD-encoding nucleic acids, and TRADD-specific binding reagents. These compositions find use in diagnostic and therapeutic methods for disease associated with undesirable cell growth, migration, differentiation and/or cytokine signal responsiveness and methods and compositions for identifying lead compounds and pharmacological agents.

L11 ANSWER 177 OF 184 USPATFULL

AN 96:75290 USPATFULL

TI Helicobacter ***pylori*** bacterial derived factor

IN Anton, Peter A., West Hollywood, CA, United States Reeve, Jr., Joseph R., Oakhurst, CA, United States Walsh, John H., Los Angeles, CA, United States Faull, Kym F., Los Angeles, CA, United States

PA The Regents of the University of California, Oakland, CA, United States (U.S. corporation)

PI US 5547844 19960820

AI US 1995-395495 19950223 (8)

DT Utility

EXNAM Primary Examiner: Housel, James C.; Assistant Examiner: Portner, Ginny Allen

LREP Rowland, Bertram I.

CLMN Number of Claims: 7

ECL Exemplary Claim: 1 DRWN No Drawings LN.CNT 507

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Chemotactin, diethyl phthalate, is shown to be a chemoattractant secreted by H. ***pylori*** Chemotactin attracts phagocytic cells with a resulting inflammatory episode. Chemotactin and its metabolites may be used for diagnosis and monitoring courses of infection by H. ***pylori*** or other chemotactin secreting organisms. In addition, chemotactin may be used in research for studying the inflammatory process, for identifying new drugs for modulating chemoattraction and activation of phagocytic cells, and for inducing an inflammatory response as a therapeutic intervention.

L11 ANSWER 178 OF 184 USPATFULL

AN 96:53195 USPATFULL

TI CagB and CagC genes of helicobacter ***pylori*** and related compositions

IN Blaser, Martin J., Nashville, TN, United States Tummuru, Murali K. R., Nashville, TN, United States Sharma, Smita A., Nashville, TN, United States

PA Vanderbilt University, Nashville, TN, United States (U.S. corporation)

PI US 5527678 19960618

AI US 1994-327494 19941021 (8)

DT Utility

EXNAM Primary Examiner: Jones, W. Gary; Assistant Examiner: Rees, Dianne

LREP Needle & Rosenberg CLMN Number of Claims: 14 ECL Exemplary Claim: 1

DOLL DACINPILLY CIGIN. 1

DRWN 2 Drawing Figure(s); 2 Drawing Page(s)

LN.CNT 1854

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A cagB gene of H. ***pylori*** is provided. This nucleic acid can be the nucleic acid consisting of nucleotides 193 through 1158 in the sequence set forth as SEQ ID NO:1, which is an example of a native coding sequence for CagB. This nucleic acid can also be in a vector suitable for expressing a polypeptide encoded by the nucleic acid. A cagC gene of H. ***pylori*** is provided. This nucleic acid can be the isolated nucleic acid consisting of nucleotides 1170 through 3830 in the sequence set forth as SEQ ID NO:3, which is an example of a native coding sequence for CagC. This nucleic acid can also be in a vector suitable for expressing a polypeptide encoded by the nucleic acid. Isolated nucleic acids that specifically hybridize with cagB and cagC are provided. CagB and CagC are associated with peptic ulceration and other clinical syndromes in humans infected with strains of H. ***pylori*** that express it.

L11 ANSWER 179 OF 184 USPATFULL

AN 95:101215 USPATFULL

TI Receptor conjugates for targeting penicillin antibiotics to bacteria

IN Krivan, Howard C., Derwood, MD, United States Blomberg, A. Lennart I., Lund, Sweden

PA MicroCarb, Inc., Gaithersburg, MD, United States (U.S. corporation)

PI US 5466681 19951114

AI US 1994-180397 19940112 (8)

RLI Continuation of Ser. No. US 1990-484568, filed on 23 Feb 1990, now abandoned

DT Utility

EXNAM Primary Examiner: Walsh, Stephen G.; Assistant Examiner: Kemmerer, Elizabeth C.

LREP Pennie & Edmonds
CLMN Number of Claims: 10

ECL Exemplary Claim: 1

DRWN 2 Drawing Figure(s); 2 Drawing Page(s)

LN.CNT 719

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A variety of conjugates useful for the treatment of infections due to pathogenic microorganisms are provided. The conjugates comprise at least one agent coupled to a receptor which binds a microorganism. Suitable agents include anti-infectives, such as antibiotics and synthetic drugs. The present invention also provides methods for treating infections in warm-blooded animals due to pathogenic microorganisms.

L11 ANSWER 180 OF 184 USPATFULL

AN 95:92690 USPATFULL

TI Campylobacter ***pylori*** antigens and uses thereof for detection of Campylobacter ***pylori*** infection

IN Blaser, Martin J., New York, NY, United States Perez-Perez, Guillermo I., Denver, CO, United States

PA Enteric Research Laboratories, Inc., Denver, CO, United States (U.S. corporation)

PI US 5459041 19951017

AI US 1988-158003 19880218 (7)

DT Utility

EXNAM Primary Examiner: Spiegel, Carol A.

LREP Ostrolenk, Faber, Gerb & Soffen

CLMN Number of Claims: 30

ECL Exemplary Claim: 1

DRWN 9 Drawing Figure(s); 9 Drawing Page(s)

LN.CNT 1279

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Antigenic compositions are disclosed for use in diagnostic kits and the like for detecting the presence of ***antibodies*** specific for Campylobacter ***pylori***, bacteria often associated with the occurrence of Type B ***gastritis*** and peptic ulcer disease. Samples of bodily fluids, for instance, may be contacted with immobilized antigen which is then washed and tested for the occurrence of significant levels of antigen/ ***antibody*** complex. Levels exceeding a predetermined positive threshold are indicative of ***antibodies*** to Campylobacter ***pylori*** in the sample tested. Kits employing the antigenic compositions of the invention preferably include means for detecting the antigen/ ***antibody*** complex such as materials and reagents for conducting an enzyme-linked immunosorbent assay, Western blot technique, liposome-based assay or other known detection tests.

L11 ANSWER 181 OF 184 USPATFULL

AN 95:47611 USPATFULL

TI Rapid in vitro test for helicobacter ***pylori*** using saliva

IN Cripps, Allan, East Maitland, Australia

Witt, Campbell, Bicton, Australia Clancy, Robert L., New Lambton, Australia

Stiel, Daniel, East Lindfield, Australia

PA Auspharm International Ltd., New South Wales, Australia (non-U.S. corporation)

PI US 5420014 19950530

AI US 1992-876524 19920430 (7)

DT Utility

EXNAM Primary Examiner: Bidwell, Carol E.

LREP Scully, Scott, Murphy & Presser

CLMN Number of Claims: 17

ECL Exemplary Claim: 1

DRWN 3 Drawing Figure(s); 3 Drawing Page(s)

LN.CNT 638

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention contemplates a method for detecting contempory infection by H. ***pylori*** in a mammal comprising contacting a mucous secretion from said mammal with an antigen component from H. ***pylori*** for a time and under conditions sufficient for an IgG ***antibody*** in said mucous secretion specific to said antigen component to form a complex therewith and then subjecting said complex to a detecting means. Preferably, the antigen component is immobilized onto a solid support.

L11 ANSWER 182 OF 184 USPATFULL

AN 95:29722 USPATFULL

TI Taga gene and methods for detecting predisposition to peptic ulceration

IN Cover, Timothy L., Nashville, TN, United States Tummuru, Murali K. R., Nashville, TN, United States Blaser, Martin J., Nashville, TN, United States

PA Vanderbilt University, Nashville, TN, United States (U.S. corporation)

PI · US 5403924 19950404

AI US 1993-53614 19930426 (8)

RLI Continuation-in-part of Ser. No. US 1992-959940, filed on 13 Oct 1992, now abandoned

DT Utility

EXNAM Primary Examiner: Parr, Margaret; Assistant Examiner: Campbell, Eggerton

LREP Needle & Rosenberg

CLMN Number of Claims: 6

ECL Exemplary Claim: 1

DRWN 3 Drawing Figure(s); 3 Drawing Page(s)

LN.CNT 1854

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides an isolated nucleic acid encoding an approximately 120-128 kilodalton antigen of Helicobacter ***pylori***, or an antigenic fragment thereof, wherein the antigen is associated with peptic ulceration. The present invention also provides methods of detecting the presence of a Helicobacter ***pylori*** strain possessing the 120-128 kilodalton antigen in a subject, comprising the steps of contacting an ***antibody*** -containing sample from the subject with a detectable amount of the tagA antigen or antigenic fragment of the present invention and detecting the reaction of the antigen or fragment and the ***antibody*** A mutant H.

pylori not expressing a functional tagA antigen is also provided.

L11 ANSWER 183 OF 184 USPATFULL

AN 93:93550 USPATFULL

TI Method and product for the treatment of ***gastric*** disease

IN Cordle, Christopher T., Centerburg, OH, United States Schaller, Joseph P., Columbus, OH, United States

PA Abbott Laboratories, Abbott Park, IL, United States (U.S. corporation)

PI US 5260057 19931109

AI US 1992-999233 19921231 (7)

RLI Division of Ser. No. US 1992-926181, filed on 7 Aug 1992 which is a continuation of Ser. No. US 1990-559793, filed on 30 Jul 1990, now abandoned

DT Utility

EXNAM Primary Examiner: Chan, Y. Christina; Assistant Examiner: Loring, Susan

LREP Drayer, Lonnie R.; Nickey, Donald O.

CLMN Number of Claims: 2

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 672

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB This invention describes a product obtained from the isolation and concentration of specific immunoglobulins (***antibodies***) derived from the mammary secretions of cows immunized with Helicobacter ***pylori***. The product is useful in preparing formulations for the treatment and/or prevention of ***gastric*** diseases.

L11 ANSWER 184 OF 184 USPATFULL

AN 93:91432 USPATFULL

TI Method and product for the treatment of ***gastric*** disease

IN Cordle, Christopher T., Centerburg, OH, United States Schaller, Joseph P., Columbus, OH, United States

PA Abbott Laboratories, Abbott Park, IL, United States (U.S. corporation)

PI US 5258178 19931102

AI US 1992-926181 19920807 (7)

RLI Continuation of Ser. No. US 1990-559793, filed on 30 Jul 1990, now abandoned

DT Utility

EXNAM Primary Examiner: Lacey, David L.; Assistant Examiner: Futrovsky, Susan

LREP Drayer, Lonnie R.; Nickey, Donald O.

CLMN Number of Claims: 3

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 684

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB This invention describes a product obtained from the isolation and concentration of specific immunoglobulins (***antibodies***) derived from the mammary secretions of cows immunized with Helicobacter ***pylori*** . The product is useful in preparing formulations for the treatment and/or prevention of ***gastric*** diseases.

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SYSTEM: OS - DIALOG OneSearch
 File 155:MEDLINE(R) 1966-2000/Dec W4
         (c) format only 2000 Dialog Corporation
*File 155: First Medline 2001 update is expected towards the end of
February. For other NLM information see Help News155.
 File 5:Biosis Previews(R) 1969-2001/Feb W3
         (c) 2001 BIOSIS
 File 73:EMBASE 1974-2001/Feb W3
        (c) 2001 Elsevier Science B.V.
*File 73: For information about Explode feature please
see Help News73.
 File 94:JICST-EPlus 1985-2001/Feb W2
         (c)2001 Japan Science and Tech Corp(JST)
*File 94: There is no data missing. UDs have been adjusted to reflect
the current months data. See Help News94 for details.
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           99070725
09774848
                        to H+,K+- ATPase, serum pepsinogen A and
                pylori in relation to gastric mucosa morphology in
patients with low or low-normal concentrations of serum cobalamins.
Jul 1998
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6/6/2 (Item 2 from file: 155) 06651765 91047803

Acid and barriers. Current research and future developments for peptic ulcer therapy.
1990

6/6/3 (Item 1 from file: 5) 11244184 BIOSIS NO.: 199800025516

Helicobacter pylori associated autoantibodies recognize Lewis antigens, and peptide epitopes of gastric H+, K+- ATPase and intrinsic factor.
1997

6/6/4 (Item 1 from file: 94)
04051065 JICST ACCESSION NUMBER: 99A0425613 FILE SEGMENT: JICST-E
Physiology and pathology. 5. Gastric secretion mechanism., 1999
?logoff hold

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*** NEW Current Year Ranges Install ***

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File 155:MEDLINE(R) 1966-2000/Dec W4

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*File 155: First Medline 2001 update is expected towards the end of February. For other NLM information see Help News155.

File 5:Biosis Previews(R) 1969-2001/Feb W3

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File 73:EMBASE 1974-2001/Feb W3

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*File 73: For information about Explode feature please see Help News73.

File 94:JICST-EPlus 1985-2001/Feb W2

(c) 2001 Japan Science and Tech Corp(JST)

*File 94: There is no data missing. UDs have been adjusted to reflect the current months data. See Help News94 for details.

Set Items Description

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6/9/2 (Item 2 from file: 155)

DIALOG(R) File 155: MEDLINE(R)

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06651765 91047803

Acid and barriers. Current research and future developments for peptic ulcer therapy.

Rademaker JW; Hunt RH

Division of Gastroenterology, McMaster University Medical Centre, Hamilton, Ontario, Canada.

Scandinavian journal of gastroenterology. Supplement (NORWAY) 1990, 175 p19-26, ISSN 0085-5928 Journal Code: UCT

Languages: ENGLISH

Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL

JOURNAL ANNOUNCEMENT: 9102 Subfile: INDEX MEDICUS

Medical therapy for peptic ulcer disease has been targeted at inhibiting acid secretion based on the belief that ulcers occur due to an imbalance between aggressive and protective factors. New antisecretory agents are unlikely to show any dramatic improvement over the success and safety of histamine H2 receptor antagonists or the recently introduced H+K+ATPase proton pump antagonist omeprazole. The development of specific muscarinic M3 and gastrin receptor antagonists will provide useful agents to suppress acid and pepsinogen secretion by alternative means and may prevent the associated hypergastrinaemia seen with anti-secretory therapy. Enhancement of mucosal defence by site protective agents will be based on a better understanding of the vascular and immune factors involved in maintaining mucosal integrity and the growth factors that regulate wound healing. Molecular techniques are likely to produce the 'model anti-ulcer' agent which will effectively inhibit acid secretion and also enhance wound healing thus providing a cure for this chronic disease. (67 Refs.)

Tags: Human

Descriptors: *Antacids--Therapeutic Use--TU; *Anti-Ulcer Agents --Therapeutic Use--TU; *Peptic Ulcer--Drug Therapy--DT; Gastric Mucosa --Physiology--PH; Helicobacter pylori; Helicobacter Infections --Complications--CO; Histamine H2 Antagonists--Therapeutic Use--TU; Intestinal Mucosa--Physiology--PH; Peptic Ulcer--Etiology--ET; Wound Healing--Physiology--PH

CAS Registry No.: 0 (Antacids); 0 (Anti-Ulcer Agents); 0 (Histamine H2 Antagonists)

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(Item 1 from file: 5)
 6/9/3
DIALOG(R)File
                5:Biosis Previews(R)
(c) 2001 BIOSIS. All rts. reserv.
           BIOSIS NO.: 199800025516
                pylori associated autoantibodies recognize Lewis antigens,
 and peptide epitopes of gastric H+, K+- ATPase and intrinsic factor.
AUTHOR: Appelmelk B J(a); Straver S(a); Claeys D; Faller G; Kirchner T;
  Negrini R; Krakowka S; Eaton K; Vandenbroucke-Grauls C M J E(a)
AUTHOR ADDRESS: (a) Vrije Univ., Amsterdam**Netherlands
JOURNAL: Gut 41 (SUPPL. 1):pA17 1997
CONFERENCE/MEETING: European Helicobacter Pylori Study Group Xth
International Workshop on Gastroduodenal Pathology and Helicobacter Pylori
 Lisbon, Portugal September 11-14, 1997
SPONSOR: European Helicobacter pylori Study Group
ISSN: 0017-5749
RECORD TYPE: Citation
LANGUAGE: English
REGISTRY NUMBERS: 9000-83-3: ATPASE ; 9001-10-9: PEPSINOGEN
DESCRIPTORS:
  MAJOR CONCEPTS: Clinical Immunology (Human Medicine, Medical Sciences);
    Gastroenterology (Human Medicine, Medical Sciences); Infection
  BIOSYSTEMATIC NAMES: Aerobic Helical or Vibrioid Gram-Negatives--
    Eubacteria, Bacteria, Microorganisms; Hominidae--Primates, Mammalia,
    Vertebrata, Chordata, Animalia; Suidae--Artiodactyla, Mammalia,
    Vertebrata, Chordata, Animalia
  ORGANISMS: human (Hominidae) -- patient; pig (Suidae);
                                                          Helicobacter -
           (Aerobic Helical or Vibrioid Gram-Negatives)
  BIOSYSTEMATIC CLASSIFICATION (SUPER TAXA): Animals; Artiodactyls;
    Bacteria; Chordates; Eubacteria; Humans; Mammals; Microorganisms;
    Nonhuman Mammals; Nonhuman Vertebrates; Primates; Vertebrates
  DISEASES: gastric atrophy--digestive system disease; type B gastritis--
    digestive system disease; Helicobacter -pylori infection--bacterial
    disease, digestive system disease, pathophysiology, pathogenesis
  CHEMICALS & BIOCHEMICALS: gastric intrinsic factor; gastric proton,
    potassium ion-ATPase --peptide epitopes; pepsinogen; Helicobacter
    -pylori -associated autoantibodies--Lewis antigen recognition
  MISCELLANEOUS TERMS:
                        Meeting Abstract
CONCEPT CODES:
  34504
          Immunology and Immunochemistry-Bacterial, Viral and Fungal
  02508
          Cytology and Cytochemistry-Human
  10064
          Biochemical Studies-Proteins, Peptides and Amino Acids
  10068
          Biochemical Studies-Carbohydrates
  10506
          Biophysics-Molecular Properties and Macromolecules
  10508
          Biophysics-Membrane Phenomena
  10806
          Enzymes-Chemical and Physical
          Enzymes-Physiological Studies
  10808
  14006
          Digestive System-Pathology
  15002
          Blood, Blood-Forming Organs and Body Fluids-Blood and Lymph
  34506
          Immunology and Immunochemistry-Immunohematology, Blood Groups
  34508
          Immunology and Immunochemistry-Immunopathology, Tissue Immunology
  36002
          Medical and Clinical Microbiology-Bacteriology
          General Biology-Symposia, Transactions and Proceedings of
  00520
             Conferences, Congresses, Review Annuals
BIOSYSTEMATIC CODES:
  06210
          Aerobic Helical or Vibrioid Gram-Negatives (1992-)
  85740
          Suidae
  86215
          Hominidae
 6/9/4
           (Item 1 from file: 94)
DIALOG(R) File 94: JICST-EPlus
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           JICST ACCESSION NUMBER: 99A0425613 FILE SEGMENT: JICST-E
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Physiology and pathology. 5. Gastric secretion mechanism.

NAKAMURA MASAHIKO (1); KISHIKAWA HIROSHI (2); ISHII HIROMASA (2) (1) Tokyo Denryoku Hosp.; (2) Keio Univ. Annual Review Shokaki, 1999, VOL.1999, PAGE.82-85, REF.21 JOURNAL NUMBER: L1627AAR UNIVERSAL DECIMAL CLASSIFICATION: 616.3-09 591.132.2.05+591.433 LANGUAGE: Japanese COUNTRY OF PUBLICATION: Japan DOCUMENT TYPE: Review ARTICLE TYPE: Review article MEDIA TYPE: Printed Publication DESCRIPTORS: gastric juice secretion; smooth muscle cell; ATPase; cotransport; ion transport; nitrogen monoxide; endothelin; pylori ; growth factor; parietal cell; pepsinogen ; Helicobacter gastric mucosa; fundus ventriculi; defence mechanism BROADER DESCRIPTORS: external secretion; secretion(physiology); digestive system physiology; cell(cytology); nuclease; hydrolase; enzyme; pyrophosphatase; biological transport; transportation; nitrogen oxide; oxide; chalcogenide; oxygen group element compound; oxygen compound; nitrogen compound; nitrogen group element compound; bioactive peptide; peptide; Helicobacter; spiral and curved bacteria; bacterium; microorganism; bioactive factor; factor; zymogen; precursor(substance); stomach; gastrointestinal duct; digestive organ; mucosa; epithelial tissue; animal tissue; biomedical tissue; organization; histomembrane; membrane and film; mechanism

CLASSIFICATION CODE(S): GH01020E; EJ06030C

*** DIALINDEX search results display in an abbreviated *** *** format unless you enter the SET DETAIL ON command. *** ?sf allscience You have 285 files in your file list. (To see banners, use SHOW FILES command) ?s pepsinogen? Your SELECT statement is: s pepsinogen? Items File 5 2: INSPEC 1969-2001/Feb W3 2731 5: Biosis Previews(R)_1969-2001/Feb W3 7 6: NTIS 1964-2001/Mar W1 2 8: Ei Compendex(R) 1970-2001/Feb W1 3 9: Business & Industry(R) Jul/1994-2001/Feb 22 227 10: AGRICOLA 70-2001/Feb 15: ABI/Inform(R) 1971-2001/Feb 22 12 16: Gale Group PROMT(R) 1990-2001/Feb 22 19: CHEM.INDUSTRY NOTES 1974-2001/ISS 200108 3 20: World Reporter 1997-2001/Feb 23 28: Oceanic Abst. 1964-2001/Mar 34: SciSearch(R) Cited Ref Sci 1990-2001/Feb W4 1517 52 35: Dissertation Abstracts Online 1861-2000/Dec 41: Pollution Abs 1970-2001/Mar 3 42: PHARMACEUTICAL NEWS INDEX 1974-2001/Feb W1 3 44: Aquatic Sci&Fish Abs 1978-2001/Feb 29 16 47: Gale Group Magazine DB(TM) 1959-2001/Feb 22 48: SPORTDiscus 1962-2001/Feb 636 50: CAB Abstracts_1972-2001/Jan 11 51: Food Sci.&Tech.Abs 1969-2001/Apr W4 53: FOODLINE(R): Food Science & 6 Technology 1972-2001/Feb 21 72 65: Inside Conferences 1993-2001/Feb W3 68: ENV.BIB. 1974-2000/NOV 1 6 70: SEDBASE 1996/Jan Q1 315 71: ELSEVIER BIOBASE 1994-2001/Feb W4 73: EMBASE 1974-2001/Feb W3 1653 74: Int.Pharm.Abs. 1970-2001/Jan 76: Life Sciences Collection_1982-2001/Dec 248 77: Conference Papers Index $\overline{1}973-2001/Jan$ 72 79: Foods Adlibra (TM) $1974-\overline{2001}$ /Feb 91: MANTIS(TM)_1880-2000/Apr Examined 50 files

777 94: JICST-EPlus 1985-2001/Feb W2 98: General Sci Abs/Full-Text 1984-2001/Jan 34 103: Energy SciTec 1974-2001/Feb B1 6 107: Adis R&D Insight_1986-2001/Feb W4 6 108: AEROSPACE DBASE_1962-2001/FEB 4 109: Nuclear Sci. Abs. 1948-1976 2 124: CLAIMS(R)/REFERENCE_2000/Q3 128: PHARMAPROJECTS 1980-2001/Feb W3 2 1 129: PHIND(Archival) 1980-2001/Feb W3 68 143: Biol. & Agric. Index 1983-2001/Jan 903 144: Pascal 1973-2001/Feb W2 29 148: Gale Group Trade & Industry DB 1976-2001/Feb 21 149: TGG Health&Wellness DB(SM) 1976-2001/Feb W2 63 409 151: HealthSTAR 1975-2000/Dec 155: MEDLINE(R) 1966-2000/Dec W4 156: Toxline(R) 1965-2000/Dec 2281 335 161: Occ.Saf.& Hth. 1973-1998/Q3 14 162: CAB HEALTH 1983-2001/Jan 117 172: EMBASE Alert 2001/Feb W3 11 174: Pharm-line(R) 1978-2001/Feb W1 10

180: Federal Register 1985-2001/Feb 22

185: Zoological Record Online(R) 1978-2001/Jan

29

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                    292: GEOBASE (TM) 1980-2001/Feb
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                    305: Analytical Abstracts_1980-2001/Feb W2 315: ChemEng & Biotec Abs_1970-2000/Dec
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                    319: Chem Bus NewsBase 1984-2001/Feb 23
                    322: Polymer Online_
       Examined 150 files
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                    342: Derwent Patents Citation Indx 1978-00/200107
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               1
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               2
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       Examined 250 files
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                    653: US Pat.Fulltext_1980-1989
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                    654: US Pat.Full. 1990-2001/Feb 20
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                    764: BCC Market Research_1989-2001/Jan
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Temp SearchSave "TD245" stored
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73: EMBASE 1974-2001/Feb W3

?rf

Ref

N1

N2

И3

2281

1653

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Ν7
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N11
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N12
             335
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N13
             315
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N15
             227
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N16
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N17
N18
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                   349: PCT Fulltext 1983-2001/UB=20010215, UT=20010201
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N20
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N23
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                   143: Biol. & Agric. Index 1983-2001/Jan
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              63
                   149: TGG Health&Wellness DB(SM) 1976-2001/Feb W2
N25
              52
                    35: Dissertation Abstracts Online 1861-2000/Dec
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                   103: Energy SciTec_1974-2001/Feb B1
              32
N27
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              31
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N36
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N37 23 305: Analytical Abstracts 1980-2001/Feb W2 N38 20 444: New England Journal of Med. 1985-2001/Feb W4 N39 16 47: Gale Group Magazine DB(TM) 1959-2001/Feb 22 14 161: Occ.Saf.& Hth. $1973-1998/Q\overline{3}$ 102 files have one or more items; file list includes 285 files.

⁻ Enter P or PAGE for more -

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N41
N42
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N43
              11
                     51: Food Sci.&Tech.Abs 1969-2001/Apr W4
N44
              11
                    172: EMBASE Alert 2001/Feb W3
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N46
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                    174: Pharm-line(R) 1978-2001/Feb \overline{W}1
N47
              10
                    266: FEDRIP 2001/Feb
N48
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N49
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                    319: Chem Bus NewsBase_1984-2001/Feb 23
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N62
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                    19: CHEM.INDUSTRY NOTES 1974-2001/ISS 200108
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                    91: MANTIS(TM) 1880-2000/Apr
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                   109: Nuclear Sci. Abs. 1948-1976
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N67
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N68
N69
                   429: Adis Newsletters (Archive) 1982-2001/Dec 12
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445: IMSWorld R&D Focus 1991-2001/Jan.W4
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N79
                   124: CLAIMS(R)/REFERENCE 2000/Q3
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N80
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N82
                   229: Drug Info. 2000/Q3
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                   303: Chapman & Hall Chemical Database 1997/Apr
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N84
                   315: ChemEng & Biotec Abs 1970-2000/Dec
N85
               2
                   322: Polymer Online_
               2
N86
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                    624: McGraw-Hill Publications 1985-2001/Feb 21
N89
               2
                    649: Gale Group Newswire ASAP (TM) 2001/Feb 20
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N94
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                    371: French Patents 1961-2000/BOPI 0052
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N100
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Your last SELECT statement was:
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Ref
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N103
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N108
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                     40: Enviroline(R) 1975-2001/Feb
N109
                0
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43: Health News Daily 1990-2001/Feb 14

102 files have one or more items; file list includes 285 files.

- Enter P or PAGE for more - ?b n2 n1 n3 n9;exs

0

N110

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File 411:DIALINDEX(R)
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DIALINDEX (R)

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*** DIALINDEX search results display in an abbreviated *** *** format unless you enter the SET DETAIL ON command. *** ?sf biotech

You have 23 files in your file list. (To see banners, use SHOW FILES command) ?show files

File Name

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- 6: NTIS 1964-2001/Feb W4
- 8: Ei Compendex(R) 1970-2001/Jan W2
- 34: SciSearch(R) Cited Ref Sci 1990-2001/Feb W2
- 65: Inside Conferences $1993-20\overline{0}1/\text{Feb}$ W2
- 71: ELSEVIER BIOBASE 1994-2001/Jan W4
- 73: EMBASE 1974-2001/Feb W1
- 76: Life Sciences Collection 1982-2001/Dec
- 94: JICST-EPlus_1985-2001/Jan W4
- 98: General Sci Abs/Full-Text_1984-2001/Dec
- 99: Wilson Appl. Sci & Tech Abs_1983-2001/Dec
- 143: Biol. & Agric. Index_1983-2 $\overline{0}$ 01/Dec
- 144: Pascal 1973-2001/Feb W1
- 155: MEDLINE(R) 1966-2000/Dec W4
- 172: EMBASE Alert 2001/Feb W1
- 266: FEDRIP 2001/Jan
- 315: ChemEng & Biotec Abs_1970-2000/Dec
- 357: Derwent Biotechnology Abs 1982-2001/Apr B1
- 358: Current BioTech Abs $1983-\overline{1}999/\text{Dec}$
- 369: New Scientist 1994-2001/Jan W4
- 370: Science 1996-1999/Jul W3
- 399: CA SEARCH(R) 1967-2001/UD=13407
- 434: SciSearch(R) Cited Ref Sci 1974-1989/Dec

TP-001

Screen; lest of extensive Chronic Sustrations

Castroenterol Spn 1107; 22:131-41 Furuta, Tet al Journal of Climae Custments
2996
Vol. 20 (Supp 2)

5 107-5111

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(Item 40 from file: 73)
DIALOG(R) File 73: EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
             EMBASE No: 1996040882
 Serum pepsinogen I in childhood Helicobacter pylori gastritis: Its
relation to mucosal peptic activity
  Yahav J.; Oderda G.; Diver-Haber A.; Fradkin A.; Keller N.; Altare F.;
Ansaldi N.; Jonas A.
  Pediatric Gastroenterology Unit, Chaim Sheba Medical Center, 52621
  Tel-Hashomer Israel
  Israel Journal of Medical Sciences (ISR. J. MED. SCI.) (Israel) 1996,
  32/1 (56-59)
  CODEN: IJMDA
               ISSN: 0021-2180
  DOCUMENT TYPE: Journal; Article
  LANGUAGE: ENGLISH
                     SUMMARY LANGUAGE: ENGLISH
DRUG DESCRIPTORS:
*pepsinogen i --endogenous compound--ec
MEDICAL DESCRIPTORS:
...adult; article; child; childhood disease--etiology--et; controlled study
; enzyme activity; enzyme blood level; female; helicobacter pylori ; human
; human tissue; major clinical study; male; pepsin secretion; stomach
antrum
CAS REGISTRY NO.: 92228-49-4 (pepsinogen i ); 9001-75-6 (pepsin a)
PLEASE ENTER A COMMAND OR BE LOGGED OFF IN 5 MINUTES
? t/s5/medium, k/46, 49, 60, 64, 65, 66, 69, 70, 71
>>>'5' valid only in keyword format
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DIALOG(R) File 73: EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
             EMBASE No: 1995161422
 Decrease in basal serum pepsinogen I levels after H. pylori eradication
 DESCENSO DE LOS NIVELES BASALES DE PEPSINOGENO I SERICO TRAS LA
ERRADICACION DE H. PYLORI
  Boixeda D.; Gisbert J.P.; Vila T.; Canton R.; Redondo C.; Martin De
Argila C.; Cano A.; Garcia Plaza A.
  C/Lagasca 13,28001 Madrid Spain
  Revista Clinica Espanola ( REV. CLIN. ESP. ) (Spain)
                                                        1995, 195/4
  (214 - 219)
  CODEN: RCESA
               ISSN: 0014-2565
  DOCUMENT TYPE: Journal; Article
  LANGUAGE: SPANISH SUMMARY LANGUAGE: SPANISH; ENGLISH
DRUG DESCRIPTORS:
...*trial--ct; *clavulanic acid--drug combination--cb; *omeprazole--drug
combination--cb; *omeprazole--clinical trial--ct; *pepsinogen i
--endogenous compound--ec; *ranitidine--drug therapy--dt; *ranitidine--drug
combination--cb; *ranitidine--clinical trial...
MEDICAL DESCRIPTORS:
*helicobacter pylori ; *duodenum ulcer--drug therapy--dt; *gastritis--drug
therapy--dt; *gram negative infection--drug therapy...
...CAS REGISTRY NO.: 95510-70-6 (omeprazole); 92228-49-4 (pepsinogen i );
    66357-35-5...
            (Item 49 from file: 73)
 5/K/49
DIALOG(R) File 73: EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
06083548
             EMBASE No: 1995114035
 Diagnosis of gastric adenocarcinoma using a scoring system: Combined
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assay of serological markers of Helicobacter pylori infection, pepsinogen I
and gastrin
  Lin J.-T.; Lee W.-C.; Wu M.-S.; Wang J.-T.; Wang T.-H.; Chen C.-J.
  Department of Internal Medicine, National Taiwan University Hospital, No
  7 Chung-Shan S Rd, Sec 1, Taipei 10017 Taiwan
  Journal of Gastroenterology ( J. GASTROENTEROL. ) (Japan) 1995, 30/2
  (156-161)
  CODEN: JOGAE
                 ISSN: 0944-1174
  DOCUMENT TYPE: Journal; Article
  LANGUAGE: ENGLISH
                     SUMMARY LANGUAGE: ENGLISH
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*gastrin--endogenous compound--ec; *pepsinogen i --endogenous compound--ec
MEDICAL DESCRIPTORS:
*helicobacter pylori ; *stomach adenocarcinoma--diagnosis--di
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 5/K/60
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DIALOG(R) File 73: EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
05042431
             EMBASE No: 1992182647
 Twenty-four-hour hyperpepsinogenaemia in Helicobacter pylori-positive
subjects is abolished by eradication of the infection
  Fraser A.G.; Prewett E.J.; Pounder R.E.; Samloff I.M.
  University Department of Medicine, Royal Free Hospital, School of
  Medicine, Rowland Hill Street, London NW3 2PF United Kingdom
  Alimentary Pharmacology and Therapeutics ( ALIMENT. PHARMACOL. THER. ) (
  United Kingdom) 1992, 6/3 (389-394)
  CODEN: APTHE
                 ISSN: 0269-2813
  DOCUMENT TYPE: Journal; Article
  LANGUAGE: ENGLISH
                     SUMMARY LANGUAGE: ENGLISH
DRUG DESCRIPTORS:
...*therapy--dt; *bismuth derivative--drug combination--cb; *metronidazole
--drug therapy--dt; *metronidazole--drug combination--cb; *pepsinogen i
--endogenous compound--ec; *pepsinogen ii--endogenous compound--ec
MEDICAL DESCRIPTORS:
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(c) 2001 Elsevier Science B.V. All rts. reserv.
             EMBASE No: 1992083613
Relationship of Helicobacter pylori to serum pepsinogens in an
asymptomatic Japanese population
  Asaka M.; Kimura T.; Kudo M.; Takeda H.; Mitani S.; Miyazaki T.; Miki K.;
  Veterans Affairs Medical Ctr., 2002 Holcombe Boulevard, Houston, TX 77030
 United States
 (Castroenterology (Castroenterology ) (United States)
                                                         1992,
  (760-766)
  CODEN: GASTA
                 ISSN: 0016-5085
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                      SUMMARY LANGUAGE: ENGLISH
DRUG DESCRIPTORS:
*biological marker--endogenous compound--ec; *immunoglobulin g antibody
--endogenous compound--ec; *pepsinogen i --endogenous compound--ec; *
pepsinogen ii--endogenous compound--ec
MEDICAL DESCRIPTORS:
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*atrophic gastritis--etiology--et; *helicobacter pylori ; *gastritis
--etiology--et
CAS REGISTRY NO.: 92228-49-4 (pepsinogen i ); 61536-72-9...
            (Item 65 from file: 73)
 5/K/65
DIALOG(R) File 73: EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
04930852
             EMBASE No: 1992071068
 Effect of helicobacter pylori on serum pepsinogen I and plasma gastrin in -
duodenal ulcer patients
  Chittajallu R.S.; Dorrian C.A.; Ardill J.E.S.; McColl K.E.L.
  Department of Medicine, Western Infirmary, Glasgow G11-6NT United Kingdom
  Scandinavian Journal of Gastroenterology ( SCAND. J. GASTROENTEROL. ) (
  Norway) 1992, 27/1 (20-24)
  CODEN: SJGRA ISSN: 0036-5521
  DOCUMENT TYPE: Journal; Article
  LANGUAGE: ENGLISH
                      SUMMARY LANGUAGE: ENGLISH
DRUG DESCRIPTORS:
*gastrin--endogenous compound--ec; *pepsinogen i --endogenous compound--ec
MEDICAL DESCRIPTORS:
*helicobacter pylori ; *gastrin blood level
CAS REGISTRY NO.: 9002-76-0 (gastrin); 92228-49-4 (pepsinogen i )
 5/K/66
            (Item 66 from file: 73)
DIALOG(R) File 73: EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
             EMBASE No: 1991331128
Helicobacter pylori in dyspeptic patients: Quantitative association with
severity of gastritis, intragastric pH, and serum gastrin concentration
  Karttunen T.; Niemela S.; Lehtola J.
  Dept. of Pathology, University of Oulu, Kajaanintie 52 D,SF-90220 Oulu
  Finland
  Scandinavian Journal of Gastroenterology, Supplement ( SCAND. J.
  GASTROENTEROL. SUPPL. ) (Norway) 1991, 26/186 (124-134)
                 ISSN: 0085-5928
  CODEN: SJGSB
  DOCUMENT TYPE: Journal; Article
  LANGUAGE: ENGLISH
                     SUMMARY LANGUAGE: ENGLISH
DRUG DESCRIPTORS:
gastrin--drug concentration--cr; gastrin--endogenous compound--ec;
pepsinogen i --drug concentration--cr; pepsinogen i --endogenous compound
MEDICAL DESCRIPTORS:
*helicobacter pylori ; *disease severity; *dyspepsia; *gastrin blood level
; *gastritis; *stomach ph
CAS REGISTRY NO.: 9002-76-0 (gastrin); 92228-49-4 (pepsinogen i )
 5/K/69
            (Item 69 from file: 73)
DIALOG(R) File 73: EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
04623565
             EMBASE No: 1991117608
 Serum pepsinogen I in duodenal ulcer
 Kolster J.
 Hospital Gonzalez Plaza, Universidad de Carabobo, Carabobo Venezuela
  GEN (GEN ) (Venezuela) 1990, 44/2 (191-198)
  CODEN: GENCA
                ISSN: 0016-3503
  DOCUMENT TYPE: Journal; Review
  LANGUAGE: SPANISH
                      SUMMARY LANGUAGE: ENGLISH
```

```
DRUG DESCRIPTORS:
...*pd; *histamine h2 receptor antagonist--drug therapy--dt; *omeprazole
--pharmacology--pd; *omeprazole--drug therapy--dt; *pepsinogen i
--pharmacology--pd; *pepsinogen i --endogenous compound--ec; *ranitidine
--pharmacology--pd; *ranitidine--drug therapy--dt
MEDICAL DESCRIPTORS:
acid secretion; helicobacter pylori; genetic marker; human; review
...CAS REGISTRY NO.: 95510-70-6 (omeprazole); 92228-49-4 (pepsinogen i );
    66357-35-5...
 5/K/70
            (Item 70 from file: 73)
DIALOG(R) File 73: EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
             EMBASE No: 1989187971
04018929
 Serum pepsinogen I and IqG antibody to Campylobacter pylori in
non-specific abdominal pain in childhood
  Oderda G.; Vaira D.; Holton J.; Dowsett J.F.; Ansaldi N.
  Pediatric Gastroenterology Section, University of Torino, Torino Italy
  Gut (GUT) (United Kingdom) 1989, 30/7 (912-916)
                ISSN: 0017-5749
  CODEN: GUTTA
  DOCUMENT TYPE: Journal
  LANGUAGE: ENGLISH
                     SUMMARY LANGUAGE: ENGLISH
DRUG DESCRIPTORS:
*immunoglobulin g antibody; *pepsinogen i ; *urease
MEDICAL DESCRIPTORS:
*abdominal pain--etiology--et; *helicobacter pylori; *immunoglobulin
blood level
CAS REGISTRY NO.: 92228-49-4 (pepsinogen i ); 9002-13-5 (urease)
            (Item 71 from file: 73)
 5/K/71
DIALOG(R)File 73:EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
            EMBASE No: 1989089258
03920265
Amoxycillin plus tinidazole for campylobacter pylori gastritis in
children: Assessment by serum IgG antibody, pepsinogen I, and gastrin
  Oderda G.; Holton J.; Altare F.; Vaira D.; Ainley C.; Ansaldi N.
  Paediatric Gastroenterology Section, University of Turin, Turin Italy
  Lancet (LANCET) (United Kingdom) 1989, 1/8640 (690-692)
  CODEN: LANCA
                ISSN: 0140-6736
  DOCUMENT TYPE: Journal
  LANGUAGE: ENGLISH
                     SUMMARY LANGUAGE: ENGLISH
DRUG DESCRIPTORS:
*bacterium antibody; *gastrin; *pepsinogen i ; *amoxicillin--drug therapy
--dt; *amoxicillin--drug combination--cb; *tinidazole--drug therapy--dt; *
tinidazole--drug...
MEDICAL DESCRIPTORS:
*helicobacter pylori ; *gastritis--diagnosis--di
CAS REGISTRY NO.: 9002-76-0 (gastrin); 92228-49-4 (pepsinogen i);
```

Background: To determine the accuracy of blood tests in predicting normal gastric mucosa confirmed by histological examination of gastric biopsy specimens. Methods: In total, 207 consecutive patients referred for upper endoscopy were included. Two biopsy specimens each from the antrum and corpus were assessed histologically for the presence of Helicobacter pylori, gastritis, and atrophy. Serum samples were studied for H. pylori antibodies by enzyme immunoassay (Pyloriset EIA-G and EIA-A) and by a rapid latex agglutination test (Pyloriset Dry); pepsinogen I was measured by an immunoenzymometric assay (Gastroset PGI), gastrin by radioimmunoassay, and parietal cell antibodies by indirect immunofluorescence. Results: In 101 (49%) of 207 patients, the gastric mucosa on histologic examination was normal. In the 63 patients aged 45 years or less, H. pylori IgG serology was negative in all 47 patients with normal gastric mucosa and none had low serum pepsinogen I levels. Among 144 patients over age 45 years, 72 had negative H. pylori IgG serology. Combining the serum pepsinogen I assay with the results of H. pylori IgG serology, 12 patients with normal serology but low serum pepsinogen I were found. Thus, 60 patients, 52 of whom showed normal gastric histology, had normal IgG serology and serum pepsinogen I. In the remaining eight patients with normal blood tests, the histologic changes were very mild. Conclusions: Although negative H. pylori IgG serology alone in younger patients, and in combination with normal serum pepsinogen I levels in older patients, reliably predicted the presence of normal gastric mucosa, gastroscopy is still recommended for patients over 45 years.

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DEVICE BRAND NAME/MANUFACTURER NAME: Pyloriset EIA-G; Pyloriset EIA-A;
Pyloriset Dry; Gastroset PGI
DRUG DESCRIPTORS:
immunoglobulin G antibody; parietal cell antibody; pepsinogen I
MEDICAL DESCRIPTORS:
*gastritis--diagnosis--di; *gastritis--etiology--et; *stomach mucosa; *
serology
stomach disease--diagnosis--di; stomach disease--etiology--et; qastroscopy;
 Helicobacter pylori; stomach biopsy; histology; antibody titer; enzyme
immunoassay; radioimmunoassay; immunofluorescence; human; nonhuman; major
clinical study; article; priority journal
CAS REGISTRY NO.: 92228-49-4 (pepsinogen I )
SECTION HEADINGS:
  004 Microbiology: Bacteriology, Mycology, Parasitology and Virology
  005 General Pathology and Pathological Anatomy
  026 Immunology, Serology and Transplantation
  048 Gastroenterology
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5/9/3 (Item 3 from file: 73)
DIALOG(R)File 73:EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
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10723429 EMBASE No: 2000212521

Long-term effect of Helicobacter pylori infection on serum pepsinogens

Kikuchi S.; Kurosawa M.; Sakiyama T.; Tenjin H.; Miki K.; Wada O.; Inaba Y.

S. Kikuchi, Department of Public Health, Aichi Medical University, 21 Aza Karimata, Nagakute-cho, Aichi 480-1195 Japan
AUTHOR EMAIL: kikuchis@aichi-med-u.ac.jp
Japanese Journal of Cancer Research (JPN. J. CANCER RES.) (United Kingdom) 2000, 91/5 (471-476)
CODEN: JJCRE ISSN: 0910-5050
DOCUMENT TYPE: Journal; Article
LANGUAGE: ENGLISH SUMMARY LANGUAGE: ENGLISH
NUMBER OF REFERENCES: 21

Serum pepsinogen values are markers of gastric mucosal status and of gastric cancer risk. The effect of Helicobacter pylori infection and

gastric cancer risk. The effect of Helicobacter pylori infection and sibship size on change of serum pepsinogen values over a seven-year span was investigated. Data from 2584 subjects with phlebotomy were analyzed both in 1989 and in 1996. The subjects were classified by H. pylori serology and sibship size (1-3 vs. 4 and more). Pepsinogen I (PG I) to II (PG II) ratio in '96 minus that in '89 was defined as DeltaPG I/II and compared among the groups. DeltaPG I/II was lower and decrease of PG I/II was more frequent among H. pylori-positive subjects than among negative subjects. The difference was owing to a decrease of PG I in all subjects and owing to an increase of PG II in those not younger than 30 years in '89. In H. pylori-positive subjects, those with a larger sibship size showed lower DeltaPG I/II and higher frequency of PG I/II decline. H. pylori infection exerts a reducing effect on PG I/II during the seven-year span. The effect of H. pylori is stronger among those with a larger sibship size, who are expected to have been infected with H. pylori in childhood. Inducing atrophy of gastric mucosa, which is reflected by a decline of PG I/II, may be one of the mechanisms through which H. pylori elevates the risk of gastric cancer.

DRUG DESCRIPTORS:

*pepsinogen--endogenous compound--ec

pepsinogen I --endogenous compound--ec; pepsinogen II--endogenous compound
--ec

MEDICAL DESCRIPTORS:

*Helicobacter pylori; *Gram negative infection stomach mucosa; stomach cancer; cancer risk; phlebotomy; serology; follow up; sibling; human; male; female; major clinical study; controlled study; adult; article; priority journal

CAS REGISTRY NO.: 9001-10-9 (pepsinogen); 92228-49-4 (pepsinogen I); 61536-72-9, 95829-35-9 (pepsinogen II SECTION HEADINGS:

004 Microbiology: Bacteriology, Mycology, Parasitology and Virology

016 Cancer

048 Gastroenterology

1998:372881 CAPLUS

129:39461 DN

Diagnosis of Helicobacter pylori infection. Serum antibody, pepsinogen, and urea breath test Goto, Akira; Fujimori, Kazuya; Kaneko, Taimei; Akamatsu, Taiji 2nd Dep. Intern. Med., Shin (1000) 27(5) 262 ΤI

ΑU

CS

Nippon Naika Gakkai Zasshi (1998), 87(5), 863-867 SO CODEN: NNGAAS; ISSN: 0021-5384

Nippon Naika Gakkai PΒ

Journal; General Review DT

LΑ Japanese

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DIALOG(R) File 73: EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
             EMBASE No: 1997358708
 Antigastric autoantibodies in Helicobacter pylori infection: Implications
of histological and clinical parameters of gastritis
  Faller G.; Steininger H.; Kranzlein J.; Maul H.; Kerkau T.; Hensen J.;
Hahn E.G.; Kirchner T.
  Dr. G. Faller, Institute of Pathology, University of Erlangen-Nurnberg,
  Krankenhausstrasse 8-10, D-91054 Erlangen Germany
  Gut (GUT) (United Kingdom) 1997, 41/5 (619-623)
  CODEN: GUTTA
                 ISSN: 0017-5749
  DOCUMENT TYPE: Journal; Article
  LANGUAGE: ENGLISH
                     SUMMARY LANGUAGE: ENGLISH
  NUMBER OF REFERENCES: 38
DRUG DESCRIPTORS:
*epitope--endogenous compound--ec; *gastrin--endogenous compound--ec; *
parietal cell antibody--endogenous compound--ec; *pepsinogen i
--endogenous compound--ec; *pepsinogen ii--endogenous compound--ec
MEDICAL DESCRIPTORS:
*gastritis; *gram negative infection; *helicobacter pylori
CAS REGISTRY NO.: 9002-76-0 (gastrin); 92228-49-4 (pepsinogen i );
    61536-72-9...
 5/K/26
            (Item 26 from file: 73)
DIALOG(R) File 73: EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
06997555
             EMBASE No: 1997283757
 Changes in gastrin and serum pepsinogens in monitoring of Helicobacter
pylori response to therapy
  Perez-Paramo M.; Albillos A.; Calleja J.L.; Salas C.; Marin M.D.C.;
Marcos M.L.; Cacho G.; Escartin P.; Ortiz-Berrocal J.
  Dr. A. Albillos, Department of Gastroenterology, Clinica Puerta de
  Hierro, San Martin de Porres, 4, 28035 Madrid Spain
  Digestive Diseases and Sciences ( DIG. DIS. SCI. ) (United States);
  42/8 (1734-1740)
  CODEN: DDSCD
                 ISSN: 0163-2116
  DOCUMENT TYPE: Journal; Article
  LANGUAGE: ENGLISH
                      SUMMARY LANGUAGE: ENGLISH
  NUMBER OF REFERENCES: 22
DRUG DESCRIPTORS:
... *endogenous compound--ec; *omeprazole--drug therapy--dt; *omeprazole
--drug dose--do; *omeprazole--drug combination--cb; *pepsinogen i
--endogenous compound--ec; *pepsinogen ii--endogenous compound--ec
MEDICAL DESCRIPTORS:
...*ulcer--drug therapy--dt; *gram negative infection--etiology--et; *gram
negative infection -- drug therapy -- dt; *helicobacter pylori
...CAS REGISTRY NO.: 95510-70-6 (omeprazole); 92228-49-4 (pepsinogen i );
    61536-72-9...
            (Item 30 from file: 73)
 5/K/30
DIALOG(R)File 73:EMBASE
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(Item 24 from file: 73)

5/K/24

(c) 2001 Elsevier Science B.V. All rts. reserv.

06756211 EMBASE No: 1997037697

Serum pepsinogen I levels and acid secretion in Helicobacter pylori associated enlarged fold gastritis
Yasunaga Y.; Shinomura Y.; Kanayama S.; Miyazaki Y.; Palacios J.J.B.; Matsuzawa Y.

Dr. Y. Yasunaga, Second Department Internal Medicine, Osaka University

Bruga

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Medical School, 2-2 Yamadaoka, Suita, Osaka 565 Japan
   Italian Journal of Gastroenterology ( ITAL. J. GASTROENTEROL. ) (Italy)
 1996, 28/8 (457-461)
   CODEN: ITJGD
                  ISSN: 0392-0623
   DOCUMENT TYPE: Journal; Article
   LANGUAGE: ENGLISH SUMMARY LANGUAGE: ENGLISH
   NUMBER OF REFERENCES: 35
 DRUG DESCRIPTORS:
 *pepsinogen i --endogenous compound--ec
 MEDICAL DESCRIPTORS:
 ...level; clinical article; female; gram negative infection--drug therapy
 --dt; gram negative infection--etiology--et; helicobacter pylori ; human;
 male; stomach parietal cell
 CAS REGISTRY NO.: 92228-49-4 (pepsinogen i ); 10361-44-1...
  5/K/31
             (Item 31 from file: 73)
 DIALOG(R) File 73: EMBASE
 (c) 2001 Elsevier Science B.V. All rts. reserv.
 06738461
              EMBASE No: 1997019932
  Percentage changes in serum pepsinogens are useful as indices of
 eradication of Helicobacter pylori
   Furuta T.; Kaneko E.; Baba S.; Arai H.; Futami H.
   Dr. T. Furuta, First Department of Medicine, Hamamatsu Univ. School of
   Medicine, 3600 Handa-cho, Hamamatsu 431-31 Japan
   American Journal of Gastroenterology ( AM. J. GASTROENTEROL. ) (United
   States) 1997, 92/1 (84-88)
   CODEN: AJGAA
                ISSN: 0002-9270
   DOCUMENT TYPE: Journal; Article
   LANGUAGE: ENGLISH
                      SUMMARY LANGUAGE: ENGLISH
   NUMBER OF REFERENCES: 20
 DRUG DESCRIPTORS:
 *pepsinogen i --endogenous compound--ec; *pepsinogen ii--endogenous
 compound--ec
 MEDICAL DESCRIPTORS:
 *gram negative infection--diagnosis--di; *gram negative infection--drug
 therapy--dt; *helicobacter pylori ; *serum
 CAS REGISTRY NO.: 92228-49-4 (pepsinogen i ); 61536-72-9...
 5/K/39
             (Item 39 from file: 73)
 DIALOG(R) File 73: EMBASE
 (c) 2001 Elsevier Science B.V. All rts. reserv.
 06388260 EMBASE No: 1996047954
Verification of decreased basal and stimulated serum pepsinogen-I levels
 is a useful non-invasive method for determining the success of eradication
 therapy for Helicobacter pylori
   Gisbert J.P.; Boixeda D.; Vila T.; De Rafael L.; Redondo C.; Canton R.;
 Martin de Argila C.
   C/Lagasca 13,E-28001 Madrid Spain
   Scandinavian Journal of Gastroenterology ( SCAND. J. GASTROENTEROL. ) (
 Norway) 1996, 31/2 (103-110)
  CODEN: SJGRA
                 ISSN: 0036-5521
  DOCUMENT TYPE: Journal; Article
  LANGUAGE: ENGLISH
                      SUMMARY LANGUAGE: ENGLISH
 DRUG DESCRIPTORS:
 *metronidazole--drug therapy--dt; *metronidazole--drug combination--cb; *
pepsinogen i --endogenous compound--ec; *tetracycline--drug therapy--dt; *
tetracycline--drug combination--cb
MEDICAL DESCRIPTORS:
adult; aged; article; clinical article; clinical trial; controlled study;
```

John Jana

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drug efficacy; female; helicobacter pylori; human; human tissue; male;
priority journal; stomach biopsy
...CAS REGISTRY NO.: 443-48-1 (metronidazole); 92228-49-4 (pepsinogen i );
    23843-90-5...
 5/K/40
            (Item 40 from file: 73)
DIALOG(R) File 73: EMBASE
(c) 2001 Elsevier Science B.V. All rts. reserv.
06376091
             EMBASE No: 1996040882
 Serum pepsinogen I in childhood Helicobacter pylori gastritis: Its
relation to mucosal peptic activity
  Yahav J.; Oderda G.; Diver-Haber A.; Fradkin A.; Keller N.; Altare F.;
Ansaldi N.; Jonas A.
  Pediatric Gastroenterology Unit, Chaim Sheba Medical Center,52621
  Tel-Hashomer Israel
  Israel Journal of Medical Sciences (ISR. J. MED. SCI.) (Israel) 1996,
  32/1 (56-59)
  CODEN: IJMDA
                ISSN: 0021-2180
  DOCUMENT TYPE: Journal; Article
 LANGUAGE: ENGLISH
                     SUMMARY LANGUAGE: ENGLISH
DRUG DESCRIPTORS:
*pepsinogen i --endogenous compound--ec
MEDICAL DESCRIPTORS:
...adult; article; child; childhood disease--etiology--et; controlled study
```

; enzyme activity; enzyme blood level; female; helicobacter pylori; human ; human tissue; major clinical study; male; pepsin secretion; stomach

CAS REGISTRY NO.: 92228-49-4 (pepsinogen i); 9001-75-6 (pepsin a)

File 155:MEDLINE(R) 1966-2000/Dec W4 (c) format only 2000 Dialog Corporation 73:EMBASE 1974-2001/Feb W1 (c) 2001 Elsevier Science B.V. ?ds Set Items Description S1 23081 HELICOBACTER PYLORI S2 190 PEPSINOGEN I 71 S1 AND S2 s3 0 H, K-ATPASE S4

RD S3 (unique items)

71

?t s5/free/1-10

S5

5/6/2 (Item 2 from file: 73) 10840015 EMBASE No: 2000321262

Evaluation of blood tests to predict normal gastric mucosa $2000\,$

5/6/3 (Item 3 from file: 73) 10723429 EMBASE No: 2000212521

Long-term effect of Helicobacter pylori infection on serum pepsinogens $2000\,$

24 26 , 30 ,71 ,31 ,40 ,46 ,49 ,60 ,64 65, 66 ,69 ,70 ,71

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DIALOG(R) File 155: MEDLIN 1:31
(c) format only 2000 idalog Concoration. All rts. reserv.
10546918
          20335634
  Antibody enzymes in "Helicobacter pylori-associated infection]
  Antitela-fermenty pri Helicolacter pylori -assotsiirovannoi infektsii.
 Konorev MR
  State Medical Institute, Vitebsk, Belarus.
  Zhurnal mikrobiologii, epidemiologii, i immunobiologii (RUSSIA) Jan-Feb
      (1) p75-9, ISSN 0372-9311
                                  Journal Code: Y90
 Languages: RUSSIAN Summary Languages: ENGLISH
  Document type: JOURNAL ARTICLE ; English Abstract
12/3/2
DIALOG(R) File 15
                    100
(c) format only 2 30 Dis o \mathcal{L}_i or All rts. reserv.
10543336
          20341254
 Noninvasive tests as a sesimitate for histology in the diagnosis of
Helicobacter pylori infection.
 Hahn M; Fennerty Ma; Corless CL; Magaret N; Lieberman DA; Faigel DO
 Division of Gastroenterology, Portland VA Medical Center and Oregon
Health Sciences University, UK 97201, USA.
 Gastrointestinal endoscopy (UNITED STATES) Jul 2000, 52 (1) p20-6,
ISSN 0016-5107
               Journal Code: FH8
 Languages: ENGLISH
 Document type: CLINICAL TRIAL; JOURNAL ARTICLE
12/3/3
DIALOG(R) File 155:METIMF 'R)
(c) format only 000 . i.e. o. t. 1.
                                         r+s. reserv.
                       Programme and American
10455773
        20307965
Factors that affect results of the 130 usea breath test in Japanese
patients.
 Chen X; Haruma K; Kam da.T; ihara M; Komoto K; Yoshihara M; Sumii K;
Kajiyama G
 Gastrointestinal Unit, First Deps tment of Internal Medicine, University
School of Medicine, Historhima; Japan.
 Helicobacter (UNIT): STATES) Jun 2000, 5 (2) p98-103, ISSN 1083-4389
Journal Code: CY4
 Languages: ENGLISH
 Document type: JOURNAL ARTICLE
12/3/4
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Clal in Lat Ln. A 1 its. reserv.
10422738
          20282082
The assessment of the degree of the colonization of the gastric mucosa by
               pylori and of the gastritis and duodenitis activity in
duodenal peptic ulcer in do ferent age groups]
 Otsinka stupenia obsimen. nia slyzovoi obolonky shlunka Helicobacter
pylori ta aktyviost !istrytu i duodenitu pry vyrazkovii khvorobi
dvanadtsiatipaloi kysaky v riznykh v kovykh hrupakh.
 Soloviova HA
 Likars'ka sprava (UKRAINE)
                           Jul 1:99, (5) p41÷3, ISSN 1019-5297
Journal Code: CIU
 Languages: UKRAINIAN Summary Languages: ENGLISH
 Document type: JOURNAL ARTICLE ; English Abstract
```

12/3/5

DIALOG(R) File 155: MEDLINE (R) Company (1)

(c) format only 2000 Dialog Corporation. All rts. reserv. 10339884 20183029 Diagnosis of Helicobacter pylori infection in patients with atrophic gastritis: comparison of histology, 13C-urea breath test, and serology. Kokkola A; Rautelin H; Puclakkainen P; Sipponen P; Farkkila M; Haapiainen R; Kosunen TU Second Dept. of Surgery, Helsinki University Central Hospital, Finland. Scandinavian journal of gastroenterology (NORWAY) Feb 2000, 35 (2) p138-41, ISSN 0036-5521 Journal Code: UCS Languages: ENGLISH Document type: JOURNAL ARTICLE 12/3/6 DIALOG(R) File 155 MENTINE (R) (c) format only 2000 bial ; Corpus con. Al rts. reserv. 10271365 20102100 Evaluation of rapid antiles tests for the diagnosis of Helicobacter pylori infection. Faigel DO; Magaret N; Corless C; isberman DA; Fennerty MB Department of Medicine, Postland Medical Center, Oregon 97201, USA. American journal of gastroente.cology (UNITED STATES) Jan 2000, 95 (1) Journal Code: 3HE p72-7, ISSN 0002-9270 Languages: ENGLISH Document type: JOURNAL ARTICLE 12/3/7 DIALOG(R) File 155: MEDIINE(L) (c) formationly 2000 Diale Countain Ration A_{max} list reserv. 10 - 21.2 10229151 20072320 antibody test: better than 🦮 Fingerstick Helicobac er pylori laboratory serological tasti.g? Laine L; Knigge K; Fairel O; Mangaret N; Marquis SP; Vartan G; Fennerty Department of Medicine, USC School of Medicine, Los Angeles, California 90033, USA. American journal or gastroente ploy, (UNITED STATES) Dec 1999, 94 (12) p3464-7, ISSN 0002-9270 Journal Jode: 3HE Languages: ENGLISH Document type: JOURNAL ARTICLE 12/3/8 DIALOG(R) File 155:MEDLINE(+ 10229150 20072319 How useful is the compact of kit for an body to Helicobacter in urine (URINELISA, in c. "bal ca. sica? Miwa H; Hirose M; Kik : S; "rai T; Iwazaki R; Kobayashi O; Takei Y; Ogihara T; Sato N Department of Gastroe. . r.l. endo University, School of Medicine, Tokyo, Japan. American journal of gastroomand ogy (UNITED STATES) Dec 1999, 94 (12) p3460-3, ISSN 0002-9270 Journal Lode: 3HE Languages: ENGLISH

Document type: JOURNAL ARTICLE

bocament type: booking introdu

12/3/9

DIALOG(R) File 155: MEDLINE F

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 $B = 2^{-N} \Delta L C = \{ 1, 2, \ldots, 2^{N} \}$

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10083653 97372316 The evaluation of tissue kallikrein in Helicobacter pylori-associated gastric ulcer disease. Naidoo S; Ramsaroop R; Bhoola R; Bhoola KD Department of Experimental and Clinical Pharmacology, University of Natal Medical School, Durban, South Africa. naidoot@und.med.ac.za Immunopharmacology (NETHERLANDS): Jun 1997, 36 (2-3) p263-9, ISSN 0162-3109 Journal Code: GY3 Languages: ENGLISH Document type: JOURNAL ARTICLE 12/3/10 DIALOG(R) File 155: MEDLINE(R) (c) formationly 200 what o correct: P is reserv. 10070108 99394787 Helicobacter pylon infertin paturn of gastritis, and symptoms in erosive and nonerosive gastroes praegeal reflux disease. Manes G; Mosca S; Laccetti M; Loniello M; Balzano A Dept. of Gastroenterology, Cardarelli Tospital, Naples, Italy. Scandinavian journal of gastroenterology (NORWAY) Jul 1999, 34 (7) p658-62, ISSN 0036-5521 Journal Code: UCS Languages: ENGLISH Document type: JOURNAL ARTICLE 12/3/11 DIALOG(R) File 155: MEDLINE(R) (c) format only 2000 Dialog Composition. All rts. reserv. Pannella A; Tattarletti.L; ...onacasa T: Brunati S; Zambianchi M Servizio Analisi Microbiel giche RCCS Policlinico San Matteo, Pavia. Minerva chirurgica (ITAL Jur. 1999, 54 (6) p411-4, ISSN 0026-4733 Journal Code: N3I Languages: ITALIAN Summa y Lam aes: ENGLISH Document type: JOURNAL A.R. TCLE. It yoush Abstract 12/3/12 DIALOG(R) File 155: MEDLINE(R, (c) format only 2000 Dialog ^ ryoration. All rts. reserv. 10001850 99372707 Usefulness of sollogical to the determinations for confirming eradication of Hermina 1200 per sufficient. Marchildon P; B: :Da:. DH; C arles C; Doobay R; Passaretti N; Peacock J; Marshall EJ; Peura. Enteric Products, Inc., Sto. Boos, New York 11790, USA. Languages: ENGLISH 1 9 0 TRIAL 12/3/13 DIALOG(R) File 155: MEDLINE(R.

The state of the state of

(c) format only 2000 Dialog Corporation. All rts. reserv.

99306116

Realities of diagnosing Helicolagter, pylori infection in clinical practice: a case for non-invasive undirect methodologies. Metz DC; Furth EE; Faigel ' , Losser; JA; Alavi A; Barrett DM; Montone K

.... The state of the s

44.5 SA: 3 Department of Pathology and Laboratory Medicine, University of Pennsylvania Medical Center, Philadelphia 19104, USA. Metzda@mail.med.upenn.edu Yale journal of biology and medicine (UNITED STATES) Mar-Apr 1998, 71 (2) p81-90, ISSN 0044-0086 Journal Code: XR7 Languages: ENGLISH Document type: CLINICAL TRIAL; JOURNAL ARTICLE 12/3/14 DIALOG(R) File 155: MEDLINE(R) (c) format only 2000 Dialog Corporation. All rts. reserv. 09976043 99311132 Prevalence of Caro, And English in symptomatic and asymptomatic children with Helilah over pylosi or sotion. Elitsur Y; Neace C; Werthermer 17; 1 10 t WE Department of Pediatric, Mar hall iniversity School of Medicine, Huntington, West Virginia 25701-01 5, USA. Helicobacter (UNITED STATES) Jun 299, 4 (2) p100-5, ISSN 1083-4389 Journal Code: CY4 Languages: ENGLISH Document type: JOURNAL ARTICLE 12/3/15 DIALOG(R) File 155: MEDLINE(R) (c) format only 2000 Dialog Corporation. All rts. reserv. 09963839 99292015 Takeshima F; Murase K; Sninokaul 1; Kohho S Second Dept. of Internal Medicine, Magasaki University School of Medicine, Japan. . Scandinavian journal of gas_oenterology (NORWAY) Apr 1999, 34 (4) p346-52, ISSN 0036-5521 Journal Code: UCS Languages: ENGLISH Document type: JOURNA! ANT CLE 12/3/16 DIALOG(R)File 155:MEDLINE(R) (c) format only 2000 Dialog Sorp asson. All rts. reserv. 99219298 Evaluation of a new impunetan matographic test for Helicobacter pylori IgG antibodies in elderily valuable pations [see comments]
Shirin H; Bruch R and G; ep 1 Mage /; Reif S; Zaidel L; Geva D; Avni Y; Halpern Z Department of Gastroenterology, The E. Wolfson Medical Center, Holon, Journal of gastroenteralczy (JAPAN) Feb 1999, 34 (1) p7-10, ISSN 0944-1174 Journal Code: BWP Comment in J Gastroenterol :999 Feb; 34(1):145-6 Languages: ENGLISH Document type: JOURNAL ARTIGUE

Constitution Temporary

14 B 1

12/3/17

DIALOG(R) File 155: MEDL: INE(R)

(c) format only 2000 Dialog Corpo. tion. All rts. reserv.

09853779 99151749 Main over the part of the part of infaction rates in two contrasting gastric cancer ris. regin set in the China. China Gastric Cancer Study Group.

```
Wong BC; Lam SK; Ching CK; Hu WH; Kwok E; Ho J; Yuen ST; Gao Z; Chen JS;
Lai KC; Ong LY; Chen BW; Wang WH, Jiang XW; Hou XH; Lu JY
    University Department of Medicine, The University of Hong Kong, China.
    Journal of gastroenterology and hepatology (AUSTRALIA) Feb 1999, 14
(2) p120-5, ISSN 0815-9319
                                                        Journal Code: A6J
    Languages: ENGLISH
    Document type: JOURNAL ARTICLE
  12/3/18
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Comporation, All rts. reserv.
                                                                                1. 12
                      99165996
09827307
 CLO antibody assay—can it say ar white in endoscopy and biopsy?

Mehdi I; Qureshi H; Ahr d P; Molyst G; Alam SE
    Pakistan Medical Resparc's Could : Jinnah Postgraduate Medical Centre,
Karachi.
                   The Journal of the Pekis on Midd al Association (PAKISTAN)
1998, 48 (7) p203-5, ISSN 0050-9132 Journal Code: KGI
    Languages: ENGLISH
    Document type: JOURNAL ARTICLE
  12/3/19
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
09788103
                      99131439
 A prospective, multidisciplingry evaluation of premenopausal women with
iron-deficiency anemia [see control ] Kepczyk T; Cremins JE; Wong Let Tki M; Smith LR; McNally PR
    Department of Medicine, Lt? Lmc isnA y Midical Center, Aurora, Colorado,
                                          A few transfer of the state of 
    American journal of gastrcenterology (UNITED STATES) Jan 1999,
p109-15, ISSN 0002-92/0 Journal Code: 3HE
    Comment in Am J Gastroenteral 1999 Jun; 94(6):1715
    Languages: ENGLISH
    Document type: JOGRNAL ARTIC'...
 12/3/20
DIALOG(R) File 155:MEDL1NE(R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
                     99052819
09775492
   Autoantibodies reacting w^{(i)} \circ \psi^{(j)} = v^{(i)} \circ \psi^{(j)} ric watigens in Helicobacter pylori
 associated body gas to the restriction of the ldren.

Ierardi E; Francavi a k; F trans B; gr ni B; Francavilla A
Chair of Gastroenters of the sit of Bar , Italy.

Italian journal of test unit of Bar (ITALY)
                                                                  y and hepatology (ITALY) Oct 1998,
30 (5) p478-80, Journal Code:
   Languages: ENGLISH
Document type: JOURNAL A:
 12/3/21
DIALOG(R) File 155:ME! INE (P)
(c) format only 2000 Dialog Corp cas on. All rts. reserv.
09765988
                     99053746
      Serologic detection of Helicobacter pylori infection with
cagA-positive strains in \sim 2000, gastric cancer, and asymptomatic
  The Department of H. licing, exeminat I ffairs Medical Center (111D),
Baylor College of *tdicine, Hou. on, TX 77030, USA.
    Journal of gastroenterolo sa(JAPAN) 1998, 33 Suppl 10 pl8-21, ISSN
```

1 1 1 1 1

```
0944-1174 Journal Code: BWP
  Languages: ENGLISH
  Document type: JOURNAL ARTICLE
 12/3/22
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
           98344635
09678643
 Etiologic aspects of chronic urticaria.
  Liutu M; Kalimo K; Uksila J; Kal mo H
Department of Dermatology, M. dical Microbiology, and Pathology, University of Turku, Finland.

International journal of documents, y (U. STATES) Jul 1998, 37 (7)
p515-9, ISSN 0011-9059 200 300 Cone: GR2
  Languages: ENGLISH
  Document type: JOURNAL AR ICI
 12/3/23
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Corco, ion. All rts. reserv.
            98436530
  Blood tests in the management of Helicobacter pylori infection.
Italian Helicobacter pylogi (tulo Group.

Vaira D; Holton J; Nonegatt M; Landi F; Ricci C; Ali A; Gatta L; Farinelli S; Acciardi C; Nonegatt M; Landi F; Ricci C; Ali A; Gatta L; Farinelli S; Acciardi C; Nonegatt Miglioli M

Department of Internal Medicing, Corresity of Bologna, Italy.

Gut (ENGLAND) Jul 1998, 14 Sur 1 ps? 46, ISSN 0017-5749
Journal Code: FVT
  Languages: ENGLIS:
  Document type: JOURNAL FRIC L; REVILW; R.VIEW, TUTORIAL
                          . . . . . . . . . . . . .
 12/3/24
DIALOG(R) File 155: MEDLINE(R,
(c) format only 2000 Dialdy Corpor tion. All rts. reserv.
                              98349343
09563472
  Role of seroconversion in actifirming cure of Helicobacter pylori
  Feldman M; Cryer B; Lev E. Petterson WL.

Department of Inter: dict University of Texas Health Science
Center at Dallas, USA.

JAMA (UNITED STATES)

Journal Code: KFR

Languages: ENGLI H

Document type: . N. ICIE
 12/3/25
DIALOG(R) File 155: MEDLINE(N)
(c) format only 2000 Dialog por: on. All rts. reserv.
           98281740
09555787
 Positive result by ser, y liceles active Helicobacter pylori infection in patients with a rop. x gastritis.
  Kokkola A; Rautelin h; Puoloskaisen P; Sipponen P; Farkkila M; Haapiainen
R; Kosunen TU
  Finland.
  Journal of clinica nd analogy (Unlied STATES) Jun 1998, 36 (6)
Languages: ENGLISH
  Document type: CL. AL- TRIAL; MOURNAL ARTICLE; RANDOMIZED CONTROLLED
TRIAL
```

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3.7

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A Commence of the second

```
12/3/26
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
                        And the State
          98201914
 Evaluation of Pyloriset Scr en, a rapid whole-blood diagnostic test for
Helicobacter pylori infection.
 Oksanen A; Veijola L; Sipponer ; Schauman KO; Rautelin H
 Herttoniemi Municipal Holvita Herinki, Finland.

Journal of clinical microbi og (UNITED STATES) Apr 1998, 36 (4)
p955-7, ISSN 0095-113.
                         Jura Co. :: h3h
 Languages: ENGLISH
 Document type: JOURNEL ASSIGLE
12/3/27
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Corporate n. All rts. reserv.
09419389
         98140657
Comparison of rapid office-i sad sero by with formal laboratory-based
ELISA testing for diagnosis of Helimobacter pylori gastritis.
 Kroser JA; Faigel DO; Furth BE; Metz DC
Department of Pathology, and Laboratory Medicine, University of Pennsylvania Medical Center, Philage of a 19104, USA.
 Digestive diseases and riches (UNITED STATES) Jan 1998, 43 (1)
p103-8, ISSN 0163-2116 Journal Hode: EAD:
 Languages: ENGLISH
 Document type: JOU WAY AR.
                    A.R.
12/3/28
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
09402375
          98127765
                        pylori spacific antibody immunohistochemistry
 Anti- Helicobacter
improves the diagnostic a. wracy : Helicobacter pylori in biopsy
specimen from patients treated v the criple therapy.
 Marzio L; Angelucci C; Gro i .; rodoro MG; Di Campli E; Cellini L
 Institute of Fisiopa ole a Medica, Universita G. D'Annunzio, Chieti,
 American journal of streen ploype (UNITED STATES) Feb 1998, 93 (2)
p223-6, ISSN 0002-9270 Journal Tode, 3HI
 Languages: ENGLISH
 Document type: JOURN.
12/3/29
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Corpo ...on. All rts. reserv.
09377359
         98060415
  Use of serum-specific again globalins A and G for detection of
               pyrori in ration in puttents with chronic gastritis by
Helicobacter
immunoblot analysis.
 Karvar S; Karch H; Fros M; Bu ghaidt W; Gross U
 Clinic for Internal Meanure, e, which sity of Wurzburg, Germany.
 Journal of clinical microbiol of (UNITED STATES) Dec 1997, 35 (12)
                                   : Hfi
p3058-61, ISSN 009 --1137
                            ou.
 Languages: ENGLISH
 Document type: JOUR: L AR: N
12/3/30
DIALOG(R) File 155: MLDI NE(P.
```

. . .

```
(c) format only 2000 Dialog Comporation. All rts. reserv.
                          Called Miller House
09341731
         98056750
How should Helicobacter pylori infection be diagnosed?

Megraud F
 Megraud F
  Laboratoire de Bacteriologie, Hopital Pellegrin, Universite de Bordeaux,
  Cance.

Gastroenterology (UNITED S'. TES) Dec 1997, 113 (6 Suppl) pS93-8,
ISSN 0016-5085 Journal Code FE3
                              17.
  Languages: ENGLISH
  Document type: JOURNAL ARTIGLE; RL EW; REVIEW, TUTORIAL
12/3/31
DIALOG(R) File 155: MEDIAINF(R)
(c) format only 2000 Dialt, (apport on. All rts. reserv.
          97465294
09310734
  Comparison of Helicobacter pylori antibody levels in children in
Riyadh [see comments]
 Abdullah AM; Gad el Rab M : 😕 Aye I; al Mazyad AS; al Sanie A
  Faculty of Medicine, King Sata University, Riyadh, Saudi Arabia.
  Tropical gastroenterology (INDIA) Apr-Jun 1997, 18 (2) p65-6, ISSN
0250-636X Journal Code: WGL
  Comment in Trop Gastroentarel 1901 and Jun; 18(2):39-40
  Languages: ENGLISH
                      \mathcal{N}_{\mathcal{A}}
  Document type: JOURNAL ARCIC/Ell
12/3/32
DIALOG(R) File 155: MEDLIKT (K
(c) format only 2000 Dialo, Corporation. All rts. reserv.
                        والأعلال والأراب
09261930
         97452309
   Helicobacter pylori infection and adenocarcinoma arising in Barrett's
esophagus.
  Quddus MR; Henley JD; Sulair in RA; Palumbo TC; Gnepp DR
  Department of Pathology, " Above Island Hospital and Brown University
School of Medicine, Providence 0.:03, USA.
  Human pathology (UNITED STATES Sep 1997, 28 (9) p1007-9, ISSN
0046-8177 Journal Code: GE'.
  Languages: ENGLISH
                                    1,
  Document type: JOHRNA A :ICLE .
                                    12/3/33
DIALOG(R) File 155 ME(G)
(c) format only 2000 Martin Cathernation Art rus. reserv.
09248691
         97432600
                            detect infection with Helicobacter
 Evaluation of salivary -
  Loeb MB; Riddell RH; Jame. Hurt R; Smaill FM
  Department of Laboratory Wedi e, McMaster University Medical Centre,
Hamilton, Ontario.
  Canadian journal of Sastreen(arclogy (CANADA) Jul-Aug 1997, 11 (5)
p437-40, ISSN 0835-7500 Jc. 11.1 Game: CR9
 Languages: ENGLISH
  Document type: JOURNA
                          i ub
 12/3/34
DIALOG(R) File 155: ME' IN
(c) format only 20.0 Ma'
                         Corpor
                                     ·. . _ rts. reserv.
09219258 97363676
Detection of serum . . . . ies to Cagh and VacA and of serum neutralizing
activity for vacual army epacterin in patients, with Helicobacter pylori
```

42 to 12 1 4

```
-induced gastritis.
  Donati M; Moreno S; Storni E; Tucci A; Poli L; Mazzoni C; Varoli O;
Sambri V; Farencena A; Cevenini R
  Sezione di Microbiologia DMCSS, Policlinico S. Orsola, University of
Bologna, Italy.
  Clinical and diagnostic laboratory inmunology (UNITED STATES) Jul 1997,
  4 (4) p478-82, ISSN 1071-412X Journal Code: CB7
 Languages: ENGLISH
  Document type: JOURNAL ARTICLE
                  er og kommer skip
Storense og kap
                                market state of the se
12/3/35
DIALOG(R) File 155: MEDLINE (A.,
(c) format only 2000 Diago, Co p ration. All rts. reserv.
                      97237655
09132463
 Molecular biology in alignos s and epidemiology of Helicobacter
pylori : PCR for the detection and AP-PCR for characterization of patient
isolates.
  Schwarz E; Plum G; Mauff G; Hastern H; Eidt S; Schrappe M; Kruis W
  Institut fur Medizinische Mizzo'z ologie und Hygiene, Universitat zu Koln,
Germany.
  Zentralblatt fur Bakteric ogte (CTRMENY) Feb 1997, 285 (3) p368-78,
               Journal Code: BD7
ISSN 0934-8840
 Languages: ENGLISH
Document type: JOURNAL ARI'1C4
12/3/36
DIALOG(R) File 155: MEDLINE(A.
(c) format only 2000 Dialog Corporation. All rts. reserv.
                    mmo or aliquation is a
09126630
         97293856
                   pylori aggastritis in a child with sickle cell anemia
    Helicobacter
and recurrent abdominal pain. France is
 Kennedy L; Mahoney DH; Redel CA
  Department of Pediatrics, Texa Children's Hospital, Baylor College of
Medicine, Houston, USA.
  Journal of pediatric hemata agy/o.cology (UNITED STATES) Mar-Apr 1997,
 Languages: ENGLISH
 Document type: JOURNA: ICTF
12/3/37
DIALOG(R) File 155: LLINF (P)
(c) format only 2000 Dial property. . . All rts. reserv.
09024567 98061117
  Immunohistochemical detecti .f Helicobacter pylori in the surface
mucous gel layer and its classificance. Shimizu T; Akamatsu T; Oi \rightarrow , We suyama T
  Second Department of err dicine, Shinshu University School of
Medicine, Matsumoti, Japan
  Helicobacter (UNITE STA: 3) Dec こっと、 1 (4) p197-206、 ISSN 1083-4389
Journal Code: CY4
 Languages: ENGLISH
  Document type: JOU. N. .... E
 12/3/38
DIALOG(R) File 155 EDL: 3)
                                   f ...! rt:. reserv.
(c) format only Die. J Corpe
09004635
          96335846
 A comparison of pur li breate antigen and whole cell antigen of
 Helicobacter pylori by E.JE. telt--study on the application and serum
diagnoses of Helicobacter pyroricurease diagnostic reagent]
```

A Aries

100

```
Chen JJ; Yang ZX; Jiang XG
  Institute of Epidemiology & Microbiology, Chinese Academy of Preventive
Medicine, Beijing.
  Chung-hua liu hsing ping hsieh tsa chih (CHINA) Feb 1996, 17 (1) p44-6
  ISSN 0254-6450 Journal Code: CQC
 Languages: CHINESE Summary Languages: ENGLISH
  Document type: JOURNAL ARTICLE ; English Abstract
               10 to 20 10 11
12/3/39
DIALOG(R) File 155: MEDLIN R)
(c) format only 2000 Dialog : roomat n. All rts. reserv.
                   08971056
         97118685
 Atopic dermatitis successfully treated by eradication of Helicobacter
 pylori.
 Murakami K; Fujioka T Mishi C Nagai J; Tokieda M; Kodama R; Kubota
T; Nasu M
 Second Department of Internation. Onto Medical University, Japan.
  Journal of gastroentecology (JAPAn, Nov 1996, 31 Suppl 9 p77-82,
ISSN 0944-1174
              Journal Code: BWF
 Languages: ENGLISH
  Document type: JOURNAL ARTICAF
12/3/40
DIALOG(R)File 155:ME AND AN
(c) format only 2000 D log Corporation. All rts. reserv.
08920382 97092272
 Value of serology (ELISE) for the diagnosis of Helicobacter pylori
infection: evaluation in parterls attending endoscopy and in those with
fundic atrophic gastritis.
  Tucci A; Poli L; Donati ''; Mazzoni C; Cevenini R; Sambri V; Varoli O;
Bocus P; Ferrari A; Paparo GF; Calotti G
  Istituto di Clinica Media e Gas Joenterologia, Universita di Bologna,
  Italian journal of gastroenterology (ITALY) Sep 1996, 28 (7) p371-6,
ISSN 0392-0623 Journal Code: A9I
 Languages: ENGLISH
  Document type: JOURNAL F TICLE
 12/3/41
DIALOG(R) File 155:MEDLINE(R) ( )
(c) format only 2000 Dialog Corpor fich. All its. reserv.
                      garage of the same
08847538 97005047
 Quantitative detection of tweentory immunoglobulin A to Helicobacter
 pylori in gastric juice: attibed capture enzyme-linked immunosorbent
  Hayashi S; Sugiyama T; Hisano K; wakawa T; Kurokawa I; Yachi A; Isogai H
; Isogai E; Yokota K; Hirai T; C 'am K; Fujii N
  Department of Microbiology, Sapp so Medical University, Japan.
  Journal of clinical labor tory analysis (UNITED STATES) 1996, 10 (2)
p74-7, ISSN 0887-8013 : rmil Codu: JLF
 Languages: ENGLISH
  Document type: JOURNAI ANTICLE
 12/3/42
DIALOG(R) File 155:MEDLINE .)
(c) format only 2000 Dialog Co position. All rts. reserv.
          96379024
08843744
 Evaluation of Pyloriset, D(y)_{y \in A} a_{y} new rapid agglutination test for
 Helicobacter pylori antibody detection.
 Lozniewski A; De Korwin JD; Conroy MC; Plenat F; Weber M
```

 $\frac{N}{24} = \frac{N}{N} + 0$

Laboratoire de Bacteriologie, Mopital Central, Centre Hospitalier et Universitaire, Nancy, France. Journal of clinical microbiolog, (UNITED STATES) Jul 1996, 34 (7) p1773-5, ISSN 0095-1137 Journal Jode: HSH Languages: ENGLISH Languages: ENGLISH Document type: JOURNAL ARTICLE 3 % 12/3/43 DIALOG(R) File 155: MEDLINE(R) (c) format only 2000 Dialog Corporation. All rts. reserv. Control of the war of 96376306 [Sinusoidal modulated currents in the therapy of chronic gastroduodenitis in children] Sinusoidal'nye modulicovamie toki v terapii khronicheskogo gastroduodenita u detei. Bukanovich OV; Khan MA; Chistov: ; Sheliapina VV; Shavrov AA; Klochkov Voprosy kurortologii, fizi erapii, i lechebnoi fizicheskoi kultury (JSSIA) Mar-Apr 1996, (2) p2:-6. ISSN 0942-3787 Journal Code: XIG Languages: RUSSIAN Summary Lanc ages: ENGLISH Document type: JOURNAL ARTICLE : English Abstract 12/3/44 DIALOG(R) File 155: MEDLINE(R) (c) format only 2000 Dialog Corporation. All rts. reserv. August 1970 A 08724007 96200503 New noninvasive tests for Hedicobacter pylori gastritis. Comparison with tissue-based gold standard. Faigel DO; Childs M; Furth EEgo-Alavi A: Metz DC Pathology Department, University of Pennsylvania Medical Center, miladelphia 19104, USA. Philadelphia 19104, USA. Digestive diseases and sciences (UNITED STATES) Apr 1996, 41 (4) p740-8, ISSN 0163-2116 Journal Code: EAD Languages: ENGLISH Document type: JOURNAL FITIC 5 12/3/45 DIALOG(R) File 155: MEULING(R) (c) format only 2000 Distoy C por ion. All rts. reserv. 08660964 96145287 Long-term follow-up of Whitob over pylori serology after successful eradication. Cutler AF; Prasad VM Section of Gastroen ...ogy, Singl. Hospital, Detroit, Michigan 48235, American journal of gasta bendere only (UNITED STATES) Jan 1996, 91 (1) p85-8, ISSN 0002-927. ...c :nal C e.: 3E Languages: ENGLISH Document type: JOURN. .. 2001: ... 12/3/46 DIALOG(R) File 155:MFDL (c) format only 200 D. g. spore Al rts. reserv. 08652592 96228243

Testing for Helicoba er plor in clinical practice.
Cutler AF

. . . .

Section of Gastroenter ogy, inai Hospital, Detroit, Michigan 48235, USA.

American journal of medicine (UNITED STATES) May 20 1996, p35S-39S; discussion 19S-41S, ISBN 6002-9343 Journal Code: 3JU

 p_{i+1}

y: . · ·

Languages: ENGLISH

Document type: JOURNAL ARTICLE; REMIEW; REVIEW, TUTORIAL

12/3/47

DIALOG(R) File 155: MEDLINE (R)

(c) format only 2000 Dialog Corporation. All rts. reserv.

96022529

Variability in serum pepsinogen levels in an asymptomatic population.

Knight T; Greaves S; Wilson A; Hengels K; Newell D; Corlett M; Webb P; Forman D; Elder J

Department of Surgery, School of Postgraduate Medicine, Keele University, Staffordshire, UK.

European journal of gastroent: plose & hepatology (ENGLAND) 7 (7) p647-54, ISSN 0954-691X Your Hal Code: B9X

Languages: ENGLISH

Document type: JOURNAL ARTICAE

12/3/48

DIALOG(R) File 155: MEDLINE(R)

(c) format only 2000 Dialog Corporation. All rts. reserv.

95256123

pylori infertion, serum pepsinogen level and gastric Helicobacter cancer: a case-control study in Japan.

Fukuda H; Saito D; Hayashi S; Hisai H; Ono H; Yoshida S; Oguro Y; Noda T; Sato T; Katoh M; et al

Endoscopy Division, National Cancer Center Hospital, Tokyo.

Japanese journal of cancer research (JAPAN) Jan 1995, 86 (1) p64-71, Journal Code: ALA

ISSN 0910-5050

Languages: ENGLISH Document type: JOURNAL ARTICLE

12/3/49

DIALOG(R) File 155: MEDLING(A)

(c) format only 2000 Dialo Congernation. All rts. reserv.

961256".

Serodiagnosis of Helicipacte. ylori infection in patients with human immunodeficiency virus infection.

Nielsen H; Andersen LP

Department of Infectivus Discuses, Hvidovre University Hospital, Copenhagen, Denmark.

APMIS (DENMARK) Dat 1.35, 103 (U) p689-92, ISSN 0903-4641 Journal Code: AMS

Languages: ENGLIS!

Document type: JOUR. L / TOE

12/3/50

DIALOG(R) File 155: MEDLINE(R)

(c) format only 2000 Dialog Compared to All ris. reserv.

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Current status sucer pyr to in reptic ulcer disease.

Falk GW

Department of Gas - 100y, leveland Clinic Foundation, OH 44195,

Cleveland Clinic journ. L. C. Badio Le (UNITED STATES) Mar-Apr 1995, 62 (2) p95-104, ISSN 0891-1150 or that Code: DBN

A MARKET CO. 1

Languages: ENGLISE - 19 - 19

Document type: JOU: IAL AR1 LE; HEW; REVIEW, TUTORIAL

```
(c) format only 2000 Dialog Corporation. All rts. reserv.
08559844
           9637.1177
 Serum anti- Helicobacter pylori antibolies and gastritis.
  Yamamoto I; Fukuda Y; Miz:ta T; Fukada M; Nishigami T; Shimoyama T
  Department of Internal Medicine, Llyogo College of Medicine, Japan.
  Journal of clinical gastroenterology (UNITED STATES) 1995, 21 Suppl 1
pS164-8, ISSN 0192-0790 Journal Code: IBG
  Languages: ENGLISH
  Document type: JOURNAL ARTICLE
 12/3/52
DIALOG(R) File 155: MEDLINE (R;
(c) format only 2000 Dialog Copper tion. All rts. reserv.
08559838
         96371171
  Endoscopic diagnosis of castrit's in relation to Helicobacter
 pylori infection and subjective symptoms.
  Department of Endoscopy, Tolyo M tro plican Komagome Hospital, Japan.
  Journal of clinical gastroe ter logy (UNITED STATES) 1995, 21 Suppl 1
pS135-9, ISSN 0192-0790 Journal Code: IBG.
  Languages: ENGLISH
  Document type: JOURNAL ARTICLE
. 12/3/53
DIALOG(R)File 155:MEDLINE(R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
         96401802
  Detection of Helicobacter pylori DNA in gastric juice by the
polymerase chain reaction: comparison with findings in bacterial culture
and the detection of the sur Ig' and serum IgG antibodies against
 Helicobacter pylori.
  Matsukura N; Onda II Toku aga A. Kano S; Yanashita K; Ohbayashi M
  First Department of surgery, No por Medical School, Tokyo, Japan.

Journal of gast interiors JAPA: Dec 1995, 30 (6) p689-95, ISSN
0944-1174
          Journal Lode: 5Wk
  Languages: ENGLISH
  Document type: JOURNAL ARTICLE
 12/3/54
DIALOG(R) File 155: ML DLIN!
(c) format only 2000 Dia a rpc. atic . All rts. reserv.
08531600
         96356566
  The value of Helicob ter major IgG antibody in estimating the
severity of gastritis : n + ld: --
  Sim JG; Kim CJ; Seo J'
  ed: . re : . (P : .A Oct 1995, 10 (5) p329-33,
  Journal of Kore
ISSN 1011-8934
                    ile ... : A!:
  Languages: ENGLISi.
  Document type: JOURNAL ANTICLE
 12/3/55
DIALOG(R) File 155: MEDI 'NF ',
(c) format only 2000 al : Co.p ation. All rts. reserv.
. 08490683
           96093505
  Use of the Clotest and 1 prise: in the identification of Helicobacter
 pylori in asymptomatic hearthy adults]
                      the Mark Street
```

والقارا عن الرئاحة في الو

DIALOG(R) File 155: MEDLINE(R)

```
Utilizacion del Clotest y Pyloriset en la determinacion del Helicobacter
    pylori en adultos sanos asintomaticos.
                          and the state of the state of the
    Santiago S
    Clinica Industrial, Caracas, Corpoven.
    G.E.N (VENEZUELA) · Apr-Jun 1995, 49 (2) p145-8, ISSN 0016-3503
Journal Code: FL2
    Languages: SPANISH Summary Languages: ENGLISH
    Document type: CLINICAL TRIAL; JOURNAL ARTICLE; English Abstract
                                       2^{-1}(A) \cdot 1 = k^{-\frac{1}{2}}
  12/3/56
DIALOG(R) File 155: MEDLING (R)
(c) format only 2000 Dialog Composition. All rts. reserv.
08477330 96138177
Diagnosis of Helice actes pyroii infection by specific gastric mucosal IgA and IgG pylori and relations.
    Veenendaal RA; Gotz Jr; Sofficial V; Kurban F; Bernards AT; Veselic M;
Pena AS; Lamers CB
    Department of Gastroenterslogy and lepitology, University Hospital
Leiden, The Netherlands.
    Journal of clinical pathology (ENGLAND) Nov 1995, 48 (11) p990-3,
                              Journal Code: HT3
ISSN 0021-9746
   Languages: ENGLISH
    Document type: CLINICAL TRIAL; JOURNAL ARTICLE
 12/3/57
DIALOG(R)File 155:MEDLINE(R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
                                                 y'ar 1 3 c
                                                 1.1
08375074 95390263
 Gastric syphilis: Andos ministry and histological features minicking
lymphoma.
   Long BW; Johnston JH : tze : V; : The vers RH 3rd; Haick A
    Department of Gast-oc be.t.pgy, Tise.ss_ppi Baptist Medical Center,
Jackson, USA.
                                      ٠.
American journal of gardroe Geraldy (MaiTED STATES) Sep 1995, 90 (9) p1504-7, ISSN 5002-92 1 Jf Gall Col: SET
   Languages: ENGLISE
    Document type: JOURNAL ARTICLE
  12/3/58
DIALOG(R) File 155: MEDLING U.
(c) format only 2000 D'aleg ' applies on. All rts. reserv.
08365146 95375892
 Serum antibody against Health act of Mori assayed by a new capture
                                          Para County Control
   Nakata H; Itoh H; Yokoya Y; Yorka H; Ni oka S; Miyamoto K; Kitamoto N;
Miyamoto H; Tanaka " : /
   Second Department of Internation Department Department of Internation Department of Internation Department Department of Internation Department Depart
Languages: ENGLIS):
    Document type: JOURNAL ARTICLE
  12/3/59
DIALOG(R) File 155: MEDI [N] 3)
(c) format only 2000 . . . . . All rts. reserv.
08352036
                  9535709
                                  pyl i intuition in recurrent abdominal pain in
childhood: comparison of diagno diagnotes spand therapy.

Chong SK; Lou Q; Espidac MN/2 Memorman SE; Croffie JM; Lee CH; Fitzgerald
                                       a time to the second
JF
```

and the contract of the second

. . .

```
Department of Pediatric Gastroenterology and Nutrition, James Whitcomb
Riley Hospital for Children, Indiana University School of Medicine, Indianapolis 46202-5225, USA.
     Pediatrics (UNITED STATES) Aug 1995, 96 (2 Pt 1) p211-5, ISSN
 0031-4005 Journal Code: OXV
     Languages: ENGLISH
     Document type: JOURNAL ARTICLE
                                                 Manufacture of the State of the Community of the Communit
   12/3/60
 DIALOG(R) File 155: MEDLINE(R)
 (c) format only 2000 Dialog Corporation. All rts. reserv.
08262153 95211627
Isolation and preliminary evaluation of a low-molecular-mass antigen preparation for invited and its of Helicobacter pylori
   immunoglobulin G antibodi. ...
    Andersen LP; Esperant F. Such as it skova M; Soucek A
     Department of Clin. tal Elekobiology, National University, Rigshospitalet,
Copenhagen, Denmark.
     Clinical and diagnostic laborator, immunology (UNITED STATES) Mar 1995,
     Languages: ENGLISH
     Document type: JOURNAL ALTICLE
                                                   12/3/61
 DIALOG(R) File 155: MEDLINE(R)
 (c) format only 2000 Dialog Corporation. All rts. reserv.
                     94351962
Importance of Helicepact applicitinfection and pepsinogen titer in hemodialysis and renal transpart title rations in Japan]

Kashiwagi T; Iino Y;; Sakaki Ter-shi A

2nd Department of Internal sicit, Nippon Medical School, Tokyo, Japan:
Nippon Jinzo Gakkai dish (Japan) Jul 1994, 36 (7) p853-7, ISSN
0385-2385 Journal Code: K.K.
 0385-2385 Journal Code: K'K
     Languages: JAPANESE Summary Languages: ENGLISH
     Document type: JOURNAL AP 'IC' E , English Abstract
  12/3/62
 DIALOG(R) File 155:ME ( INE R)
 (c) format only 20% alog Cartanata. Ale rts. reserv.
08226840 94269548
         Helicobacter pylch est in dyspeptic children. A long-term
follow-up after treatment with the initial beamuth subcitrate and tinidazole.
Ashorn M; Ruuska 1; Kari ski R; M Tinen-A; Maki M
Dept. of Clinical Met. c. s; Unive of Tampere, Finland.
Scandinavian journal as 30 ms ogy (NORWAY) Mar 1994, 29 (3)
p203-8, ISSN 0036-5521 des 1000 ms.
    Languages: ENGLIS!
    Document type: "Common AT AND
   12/3/63
 DIALOG(R) File 155: MEDLINE(R)
 (c) format only 2000 Dialog ...po. tion. All rts. reserv.
                     94147893
 08197851
       Helicobacter pyria magnetic for d-cobalamin malabsorption.
     Carmel R; Perez-Perez GI; E 1 MJ
Department of Medicine, & arsity of Southern California School of Medicine, Los Angeles.
Digestive diseases and scie s (UNITED STATES) Feb 1994, 39 (2) p309-14, ISSN 0163-2116 Journal idea EAD
     Contract/Grant No.: Di. 12640, DK, NIDDK; MO1 RR-43, RR, NCRR
```

The Control of the Co

4. 14.

Languages: ENGLISH

Document type: JOURNAL ARTICLE

```
12/3/64
```

DIALOG(R) File 155: MEDLINE(R)

(c) format only 2000 Dialog Corporation. All rts. reserv.

95326721

Comparative evaluation of culture techniques and ELISA test in detection

of Helicobacter pylori in.ec. ... I Porownawcza ocena posiewu testu ELISA w wykrywaniu zaka.ANG.zenia Helicobacter pylori

Sito E; Karczewska & Arczeba C & Sart Ck &; Oleksy J; Nowak-Sadzikowska J; Maj A

Klinika Chorob Wallettenyes Wojskewego Szplada Klinicznego w Krakowie. Medycyna doswiadczalna i mi wobiatogla (POL: 5) 1994, 46 (4) p301-4, ISSN 0025-8601 Journa Code: WE

Languages: POLISH Summary Languages: ENGLISH Document type: JOURNAL ARTICLE ; Emglish Fbstract

12/3/65

DIALOG(R)File 155:MEDLINE(R)

(c) format only 2000 Dialog Corporation All rts. reserv. - Corporation

08059230 95066140

Diagnosis of Helicobacter py ri infection] Diagnostik der Helicobacte. Dy i -Infektion. Meryn S

Neue Wiener Privatklinik, Wian.

Wiener klinische Wochenschrift (AUSTRIA) 1994, 106 (17) p559-62,

ISSN 0043-5325 Journal; Code: XOP page 1875

Languages: GERMAN Summary Languages: ENGLISH

Document type: JCURNAL AF'ICLF; REVIEW; REVIEW, TUTORIAL; English Abstract

12/3/66

DIALOG(R) File 155: MF: L:NE(R)

(c) format only 2000 what common ich. All rts. reserv.

08003781 94376366

Evaluation of polymera. . And reaction for diagnosis of Helicobacter - weight pylori infection]

1

Takagi A; Ohta U; shi Triksto K; Kobayashi H; Harasawa S; Miwa T; Kamiya S

Department of liter al Medicine (1984) University School of Medicine. Nippon Shokakii / Gakkai (1984) Aug 994, 91 (8) p1277-82,

ISSN 0446-6586 Journal Languages: U.NF. NESF jua EN SH Document type: JOL ..., Er Abs lact

12/3/67

DIALOG(R) File 155: MEDLINE (R)

(c) format only 2000 Dialog (c) - . . . n. All rts. reserv.

94347336

The humoral impact x_0, x_0, y_0, y_0 and sobacter pylori infection in children with recurrent abdominal years.

APMIS (DENMAGE).

APMIS (DENMARK) an 94 / 1 - 7-64, ISSN 0903-4641 ournal Code: AMS
Languages: ENGLl.i.

Journal Code: AMS

No office and

```
3 M. A. 3
 12/3/68
DIALOG(R) File 155: MEDLINE(R) (R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
                 outres relations of the
         94327747 - 38 - 148 (4.25) - U. S. A.C. C.
07974929
  Serological detection of Heilcobacter
                                           pylori antibodies in children
and their parents.
  Best LM; Veldhuyzen v n Zanten S.; Sherman PM; Bezanson GS
  Department of Microbiology Valloria General Hospital, Halifax, Nova
Scotia, Canada.
  Journal of clinical microscole (ULITED STATES) May 1994, 32 (5)
Languages: ENGLIS:
  Document type: JOURNAL ARTICAL
 12/3/69
DIALOG(R) File 155: MEDLINE(R;
(c) format only 2000 Dialog Corporat .. Al rts. reserv.
                        07885427 94172050
  Evaluation of a new immunodiagnotic assay for Helicobacter pylori
 antibody detection: correlation with histopathological and
microbiological results.
  Pronovost AD; Rose SL; Pawlak JW; Robin H: Schneider R
  Quidel Corporation, San Diego, California 92121.
  Journal of clinical microbiology (UNITED STATES) Jan 1994, 32 (1)
p46-50, ISSN 0095-1137 Jou nal Code: HSH Languages: ENGLISH
  Document type: JOURNAL ARTICLE
 12/3/70
DIALOG(R) File 155: MEDLINE(E)
(c) format only 2000 Dialog Comporation. All rts. reserv.
                                   . . :
07835595
         94268951
Evaluation of a cormercial enzymentinked immunosorbent assay (ELISA) kit
for serological diagnosis of Hel placter pylori infection in a group
of non-ulcer dyspepsia sufferess.
  Ching CK; Thompson S; Buxton C; lga e 0; Holmes GK
  Department of Medicine, perland Royal Infirmary, Derby, UK. Postgraduate medical journa (EN P. NI Jun 1993, 69 (812) p456-60,
                Journa Collect FX
ISSN 0032-5473
  Languages: ENGLISH
  Document type: JC * L ARTI J
 12/3/71
DIALOG(R) File 155: MEDL1NE (R)
(c) format only 2000 Dialog or, . wic . All its. reserv.
07778580
         93175455
 Serum 13C-bicarbonate s: s: figure f gastric Helicobacter pylori urease activity.
  Moulton-Barrett R; Friedal Lor G; Michener R; Gologorsky D
  Section of Gastroenterology, rans Affairs Medical Center, Martinez,
American journal of jartherology (UNITED STATES) Mar 1993, 88 (3) p369-74, ISSN 0002-9 J c. .al ode: 3HE. Languages: ENGLISH
  Document type: JOUENAL A GL.
```

Alternative Control of the

```
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
         94280304 by the a second of a
  Serological examinations in patients with Helicobacter pylorinfections.
infections.
 Gosciniak G; Przondo-Mordarska A; Matysiak-Budnik T; Knapik Z
 Department of Microbiology, Medical Academy, WrocLaw, Poland.
 Archivum immunologiae et therapiae experimentalis (POLAND)
                                                               1993, 41
(5-6) p309-13, ISSN 0004-069X Cournal Code: 790
 Languages: ENGLISH
Document type: CLINICAL TRIAL; JOURNAL ARTICLE
12/3/73
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 bialog orposition. All rts. reserv.
07694013
         94085142
Role of Helicobacter pyloni segretary in evaluating treatment success.
 Cutler A; Schubert A; Schubert T
 Division of Gastroenterology, Henry Ford Hospital, Detroit, Michigan.
 Digestive diseases and science. UNITED STATES) Dec 1993, 38 (12)
p2262-6, ISSN 0163-2116 Journal C.de: EAD
 Languages: ENGLISH
 Document type: JOURNAL APIL ME
12/3/74
DIALOG(R) File 155:MEDLINE(R)
(c) format only 2000 Dialog Composation All rts. reserv.
         93382585
 Costs and effectiveness of diagnosis and treatment of patients with
dyspepsia, determined with the a computer model (see comments)]
 Kosten en effectiviteit van liagnastiek en behandeling van patienten met
dyspepsie, bepaald met een con estermodel. -
 Schipper CK; Rutten FF; Loft ald R.
 Erasmus Universiteit, Insuit.u. voom Medische Technology Assessment,
Rotterdam.
 Nederlands tijdschrift voor genetskunde (NETHERLANDS) Aug 28 1993, 137
(35) p1767-71, ISSN 0028-?16 Journal Code: NUK
 Comment in Ned Tilds. For Geneeskd 1993 Oct 30:137(44):2274-5
 Languages: DUTCH Summary Languages: ENGLISH
 Document type: JOURNAL ARTICLE : English Abstract
12/3/75
DIALOG(R) File 155 ... DLI.
(c) format only 2000 Di har Component on. All rts. reserv.
07593273 93343747
 Gastric syphilis. Primary . . . s by castric biopsy: report of four
 Fyfe B; Poppiti RJ 1; Some son MJ
Arkadi M. Rywli promise to logy and Laboratory Medicine, Mount
Sinai Medical Center fit of Mia F. 33140.
 Archives of pathology & borse, y medicine (UNITED STATES) Aug 1993,
117 (8) p820-3, ISSN 0003-3985 purnal Ccde: 79Z
 Languages: ENGLISH
 Document type: JOURNAL ARPIGIE
12/3/76
DIALOG(R) File :55 ... Li (A) +
(c) format only 2000 Diago proporation, all rts. reserv.
                    1.1.1
07589200 93332059
                             1440. J.
```

Ottagrant

```
Apparent reversal of early gastric mucosal atrophy after triple therapy for Helicobacter pylori:
 Borody TJ; Andrews P; Jankiewicz E; Ferch N; Carroll M
 Centre for Digestive Diseases, NSW, Australia.
 American journal of gastroenterology (UNITED STATES) Aug 1993, 88 (8)
p1266-8, ISSN 0002-9270 Journal Code: 3HE
 Languages: ENGLISH
 Document type: JOURNAL ARTICLE
12/3/77
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Comporation. All rts. reserv.
07569772
          93301099
  Diagnosis of Helicobactes pylori infection by using pyloriset EIA-G
and EIA-A for detection of porch impunoglobulin G (IgG) and IgA antibodies
[published erratum appears 🏚 💯 Clin Microbiol 1993 Sep;31(9):2556]
 Kujari H; Gronfors R; Nurmi H; Raiha
I; Stahlberg MR; Leino R
 Orion Corporation, Orion Diagnestica, Espoo, Finland.
 Journal of clinical microbiology (UNITED STATES) Jun 1993, 31 (6)
p1450-3, ISSN 0095-1137 Journal Code: HSH
 Languages: ENGLISH
 Document type: JOURNAL ARTICLE
12/3/78
DIALOG(R) File 155: MEDLINE (R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
07541184
         93251516
 Antigens for the ELISA test for serodiagnosis of Helicobacter pylori;
infection] And the Management of the
 Antigeny pre ELISA tesir all serodiagnostiku infekcie Helicobacter
pylori .
             。                程,
 Buchvald D; Buchvaldo 1 D
 Ustav imunologie Univerzity Komenskeho, Bratislava.
 Ceskoslovenska epidemiologie, mikr biologie, imunologie (CZECHOSLOVAKIA) -
 Mar 1993, 42 (1) p16-21, ISSN 0009-0522 Journal Code: CSH
 Languages: SLOVAK Summary Languages: ENGLISH
 Document type: JOURNAL ARTICLE ; Lighish Abstract
12/3/79
DIALOG(R) File 155: MEDLINE (L)
(c) format only 2000 real Correstation. All rts. reserv.
07424520
          92165002
 Relationship of Relicobacter pylori to serum pepsinogens in an
asymptomatic Japanese populacion.
 Asaka M; Kimura T; Kudo M. Takeca T; Mitani S; Miyazaki T; Miki K; Graham
 Third Department of I ternal Medicine, Hokkaido University School of
Medicine, Sapporo, Japan.
 Gastroenterology (UNITED
                            STATES), Mar 1992, 102 (3) p760-6, ISSN
           Journal Code: FH3
 Contract/Grant No.: DK 39919, DK, NIDDK
 Languages: ENGLISH :
 Document type: JOURNAL ART . M.L.
12/3/80
DIALOG(R) File 155: MLULINE(R)
                            (c) format only 2000 Dialog Corporation. All rts. reserv.
                         . : •
07419107
          91318374
```

Serum immune response to Helicobacter pylori in children:

All British Com

.\

epidemiologic and clinical applications. De Giacomo C; Lisato L; Negrini R; Licardi G; Maggiore G Clinica Pediatrica dell'Universita di Pavia, IRCCS Policlinico S. Matteo, Journal of pediatrics (UNITED STATES) Aug 1991, 119 (2) p205-10, ISSN 0022-3476 Journal Code: JLZ Languages: ENGLISH Document type: JOURNAL ARTICLE 12/3/81 DIALOG(R) File 155: MEDLINE(R) (c) format only 2000 Dialog Co polition. All rts. reserv. 91064694 Helicobacter oyungi with gurmaic carcinoma. Serum antibody prevalence in populations with the masking cancer risks. Correa P; Fox J; Forthar, 3; RALZ 19 Lin YP; Zavala D; Taylor N; Mackinley D; de Lima E; Portilla H; st a! Department of Pathology, legislana State University Medical Center, New Orleans 70112. Cancer (UNITED STATES) Dec 15 1990, (6 (12) p2569-74, ISSN 0008-543X Journal Code: CLZ Contract/Grant No.: CA-28842, CA, NCI; AI-25590, AI, NIAID; AI25631, AI, Languages: ENGLISH Document type: JOURNAL ARTICLE DIALOG(R) File 155: MEDLINE (R) 12/3/82 (c) format only 2000 Dialog Coop lation. All rts. reserv. 07406097 93195430 / Wild & Court A Clinical significative c .i. - Halicobacter pylori antibody in the diagnosis of Helicomute: pyror infection in chronic gastritis] Negayama K; Terada S; Kawan ini ... Department of Clinical Lawrator, Kagawa Medical School. Kansenshogaku gasshi (MAN) v (1992, 66 (11) p1597-8, ISSN 0387-5911 Journal C Languages: JAPINES Document type: 'OU' - LE 12/3/83 91178057 07357471 Evaluation of an instantial as assay for specific detection of immunoglobulin G antipod assay lagainst Helicobacter pylori, and antigenic cross-reactivy Latwers ori and Campylobacter jejuni. Faulde M; Putzker dertes T, dertes T Federal Republic of Germany. Journal of clinical 19 (2) 19323-7, ISSN 0095-1187 19 (2) 196. HS Languages: ENGLISE Document type: JOHAN AP IF

12/3/84

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07336365 90197249

Helicobacter pyrolic in the children with chronic diarrhoea and malnutrition. to all or the second of the se

```
Sullivan PB; Thomas JE; Wight DG; Neale G; Eastham EJ; Corrah T;
Lloyd-Evans N; Greenwood BM
 MRC Laboratories, Fajard, The Gambia.
  Archives of disease in childhood (ENGLAND) Feb 1990, 65 (2) p189-91,
ISSN 0003-9888
               Journal Code: 6XG
  Languages: ENGLISH
  Document type: JOURNAL ARTICLE
                 7
12/3/85
DIALOG(R) File 155: MEDL1NE(R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
         93071396
07169161
  Helicobacter pylori, gratritis and ulcers in pediatrics.
  Judd RH
 University of Wiscons: Hospitally Edison.
Advances in pediatros (* TD STATES) 1992, 39 p283-306, ISSN
0065-3101 Journal Code. 200
  Languages: ENGLISH
  Document type: HISTORIC. ARTICLE; JOURNAL ARTICLE; REVIEW, REVIEW,
TUTORIAL
12/3/86
DIALOG(R) File 155: MEDLINE (R)
(c) format only 2000 Dialog Corpora on. All rts. reserv.
07145544
         93033608
  Detection of Helicobacter pylori infections by antibody etermination]
 determination]
 Nachweis von Helicobester - pylori -Infektionen
                                                                 durch
Antikorperbestimmung.
 Briedigkeit H; Montag T; Spiridonow PS; Sielaff F; Wack R; Held C; Hantke
                          tary XII, 100
  Universitatsklinik fur Innere Medizin Theodor Brugsch, Medizinischen
Fakultat, Humboldt-Universitat zu Berlin.
  Zeitschrift fur arziliche Fo. bildung (GERMANY) Sep 10 1992, 86 (17)
p869-72, ISSN 0044-2178 Jo n 1 Code: XS6
 Languages: GERMAN
  Document type: JOURNAL AR [CLE
 12/3/87
DIALOG(R)File 155:MEDLINE(F)
(c) format only 2000 Palo Corpor ion. All rts. reserv.
07084474 92351036
  Serodiagnosis of Hel Öa. r /lori-associated gastritis with a
monoclonal antibod, compe ti enz me-linked immunosorbent assay.

Negrini R; Zanella I; A; ones C; Verardi R; Ghielmi S; Albertini
A; Sangaletti O; Lazwaroni M, E unchi orro G
  Institute of Chemistry, School of Gaicine, University of Brescia, Italy.
  Scandinavian journal gas (Jenterology (NORWAY) Jul 1992, 27 (7)
p599-605, ISSN 0036-5511 Jr 121 de: UCS
  Languages: ENGLISH
  Document type: JOURNAL A.
 12/3/88
DIALOG(R) File 155. FF JE(R)
(c) format only 2000 Dial g All rts. reserv.
06878854
          92206121
 The role of serology in the diagnosis of Helicobacter (Campylobacter)
 pylori infection]
 Interets de la serolo, dara la detection de l'infection a Helicobacter
 (Campylobacter) pylori ...... :
```

1

: . iq : * * * * * . * . * . * . * .

```
Fannes F; Pierard P; Baise E; Hulin G
    Laboratoire de Recherche et Developpement, Wavre, Limal.
    Acta gastro-enterologica delgica (BELGIUM)
                                                                                         Sep-Dec 1991, 54 (5-6)
p368-74, ISSN 0001-5644 Journal Code: 0NY
    Languages: FRENCH Summary Languages: ENGLISH
    Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL; English
Abstract
                                 1.5
                                             The state of the s
  12/3/89
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialon rportion. All rts. reserv.
                                          4.
06843074
                     92105332
 Characterization of intermediate species-specific 19-kilodalton
outer membrane protein . om i : bacter pylori by using a monoclonal
   Drouet EB; Denoyel GA; Londe M; Wallano E; Andujar M; de Montolos HP
    Division of Infectious Da: es, I stitut Pasteur de Lyon, France.
    Journal of clinical microbiol 3y (UNITED STATES) Aug 1991, 29 (8)
p1620-4, ISSN 0095-1137 Journ 1 % le: HSH
    Languages: ENGLISH
    Document type: JOURNAL ARTICLE
 12/3/90
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Corporation. All rts. reserv.
                                                \gamma_{ij} = \chi_{ij}^{(n)}(z) (6)
06835128 92090747
   Long term serological salve llouce after treatment of Helicobacter
pylori infection.

Veenendaal RA; Pena AS; Me_ + JL; Endtz HP; van der Est MM; van Duijn W; Eulderink F; Kreuning J; Lamers B
    Department of Gastroenterology, Leiden University Hospital, The
Netherlands.
   Gut (ENGLAND) Nov 1991, 32 (11) p1291-4, ISSN 0017-5749
Journal Code: FVT
   Languages: ENGLISH
    Document type: JOURNAL ARTI: LE
 12/3/91
DIALOG(R) File 155: MEDI | NE(R
(c) format only 2000 D alog C oc at .... All rts. reserv.
                  92022172
       Diagnosis of gastritis a sed by Helicobacter pylori in children
Czinn SJ; Carr HS; Speck. WT 🕝
   Department of Pediatrics, School of Medicine, Case Western Reserve
University, Cleveland, Ohic.
   Reviews of infectious diseases (UNITED STATES) Jul-Aug 1991, 13 Suppl
8 pS700-3, ISSN 0162-0886 | Gaurnal Code: SXN
   Contract/Grant No.: P. 258.8, A., NIAID
   Languages: ENGLISH
   Document type: JOURNAL AFTICLE
 12/3/92
DIALOG(R) File 155:MEDI LNE( )
(c) format only 2000 Hialog Componation. All rts. reserv.
06765609
                    91367230
       Helicobacter
                                     pylori anfection and gastric carcinoma among Japanese
Americans in Hawaii [see commuts].
   Nomura A; Stemmermain GN; Chyou PH; Kato I; Perez-Perez GI; Blaser MJ
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Japan-Hawaii Cancer Study, Kuakini Medical Center, Honolulu 96817.

New England journal of medicine (UNITED STATES) Oct 17 1991, 325 (16) p1132-6, ISSN 0028-4793 Journal Code: NOW Contract/Grant No.: R01-CA-33644, CA, NCI Comment in N Engl J Med 1991 Oct 17;325(16):1170-1 Languages: ENGLISH Document type: JOURNAL ARTICLE $\mathcal{L} = \mathcal{L}$ 12/3/93 DIALOG(R) File 155: MEDLINE(R) (c) format only 2000 Dialog Corporation. All rts. reserv. ÷ • 91258575 06708633 Estimation of prevalence of elicobacter pylori infection in an asymptomatic elderly population, omparing [140] urea breath test and serology. Newell DG; Hawtin PR; Stacew Al. MadDougall MH; Ruddle AC Public Health Labor: ory Jer 10 Centre for Applied Microbiology and Research, Porton Down, Salisbary, Wilshire. Journal of clinical pathology (EMGLAND) May 1991, 44 (5) p385-7, ISSN 0021-9746 Journal Code: H73 Languages: ENGLISH Document type: JOURNAL ARTICLE 12/3/94 DIALOG(R) File 155: MEDLINE(R) (c) format only 2000 Dialog Corporation. All rts. reserv. 91257543 06708144 A novel enzyme immunessay for serediagnosis of Helicobacter pylori infection. a (Sugiyama T; Imai K; Yoshid H; Takayama Y; Yabana T; Yokota K; Oguma K; Department of Internal Medicaine, Sapporo Medical College, Japan. Gastroenterology (UNITED STATES) Jul 1991, 101 (1) p77-83, ISSN Journal Cole: Fig. 0016-5085 Languages: ENGLISH Document type: JOURNAL ART THE 12/3/95 DIALOG(R) File 155:MEL (R) (c) format only 2000 . alog orpora ion. All rts. reserv. 1. 06708124 91257508 Positive serum antibod and a qualive tissue staining for Helicobacter pylori in subjects with a copy copy gastritis [see comments] Karnes WE Jr; Samloff ; S. rai. M; Kekki M; Sipponen P; Kim SW; Walsh JH Center for Ulcer Resear of and Aucation, Veterans Administration Medical Center, Los Angeles, Califo ni Gastroenterology JNLT D ST... Jul 1991, 101 (1) p167-74, ISSN 0016-5085 Journal Cods: EH3 Contract/Grant No.: D..17328 , N. DD-Comment in Gastroentar logy : 92 Feb:102(2):744-5 Languages: ENGL *: In. Document type: 12/3/96 DIALOG(R) File 155: MEDLINE (c) format only 2000 Dialog corpora on. All rts. reserv. 479 514 06500744 91041503 Committee of the state of the Detection of antibodies to Heliupbacter pylori with the immunoenzyme test and indirect immunofluores ince]

Nachweis von Antikerpern gegen Helicobacter pylori mit Enzymimmuntest

1 44 E

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und indirekter Immunfluoreszenz.
  Abb J; Striegel K; Frunmorgen P
  Mikrobiologisches Institut, Krankenanstalten Ludwigsburg.
  Leber, Magen, Darm (GERMANY) Sep 1990, 20 (5) p224-30, ISSN 0300-8622
Journal Code: L3P
  Languages: GERMAN Summary Languages: ENGLISH
  Document type: JOURNAL ARTICLE ; English Abstract
                       . . ,
 12/3/97
DIALOG(R) File 155: MEDLINE(R)
(c) format only 2000 Dialog Corporation All rts. reserv.
          90338484
06429894
Comparison of ELTSA antigonometric eparations alone or in combination for serodiagnosing Helicobather to grain fections.

Hirschl AM; Rathbone BD; Wyatt JT: Berger J; Rotter ML
  Hygiene-Institute, University of Vienna, Austria.
  Journal of clinical pathology (ENGLAND) Jun 1990, 43 (6) p511-3,
ISSN 0021-9746
                Journal Code: HT3
  Languages: ENGLISH
  Document type: JOURNAL ARTICLE
 12/3/98
DIALOG(R) File 155: MEDLINE(R;
(c) format only 2000 Dialog Comporation. All rts. reserv.
          90282212
06406478
Endoscopy in dyspeptic patient: ; gastric mucosal biopsy useful? [see
comments]
  Vaira D; Holton J; Osborn J; OD Dana L; Romanos A; Falzon M; McNeil I
  Department of Gastroenterology, Middlesex Hospital, London, England.
  American journal of gastroenterclogy (UNITED STATES) Jun 1990, 85 (6)
p701-4, ISSN 0002-9270 Journal Code: 3HE
  Comment in Am J Gastroentercl 1991 May; 86(5):647-8
  Languages: ENGLISH
  Document type: J JRNAL ' TIT' E
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↔ File 155:MEDLINE(R) 1966-2000/Dec W4
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*File 155: First Medline 2001 update is expected towards the end of February. For other NLM information see Help News155.

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1 ATPASE, CALCIUM
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E19
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E20
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               1 ATPASE, DYNEIN
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               1 ATPASE, FO
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                 ATPASE2
E35
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                 ATPASE2A
E36
                 ATPASE4
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?s e3 or e7 or e32
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             715 ATPASE //H(+)-K(+)-EXCHANGING
            3198 ATPASES
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  E42
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           2 ATPASICAS
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           1 ATPASICO
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           18 ATPASIQUE
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  ?e e24
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1 HYDROGEN, POTASSIUM ATPASE
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               Х
  R6
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               Х
  R7
            0
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               B 11 ADENOSINETRIPHOSPHATASE
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               715 DC=D8.586.277.40.25.300.
                 O ADENOSINETRIPHOSPHATASE, HYDROGEN, POTASSIUM
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                 0 H(+)-K(+)-TRANSPORTING ATPASE
                 0 HYDROGEN, POTASSIUM ATPASE
                 0 HYDROGEN, POTASSIUM, ADENOSINETRIPHOSPHATASE
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        S3
             27423 R1-R8
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S3
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?s (s1-s5) and (helicobact? or pylori or pylor or pyloris or pyloridis or hpylori or hf
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>>>"S5" does not exist
>>>Both terms in the range must be of the same type
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elis)
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                 S2
           27423 S3
           59018
                 S4
           11767
                 HELICOBACT?
           12258 PYLORI
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                 PYLORIS
             175
                 PYLORIDIS
               0
                 HPYLORI
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                 HFELIS
      S5
             133
                  (S1-S4) AND (HELICOBACT? OR PYLORI OR PYLOR OR PYLORIS OR
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Ref
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E3
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E4
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                  PEPSINOGEN (1-12)
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               7 PEPSINOGEN A
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                  PEPSINOGEN A --BIOSYNTHESIS --BI
E9
         56
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E10
         12
                  PEPSINOGEN A -- CHEMISTRY -- CH
E11
         3
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                  PEPSINOGEN A --GENETICS --GE
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         1
E23
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                  PEPSINOGEN C --BLOOD --BL
                  PEPSINOGEN C -- CHEMISTRY -- CH
E24
          4
          Enter P or PAGE for more
?s e5-e17
             126 PEPSINOGEN A
               9 PEPSINOGEN A --ANALYSIS --AN
               1 PEPSINOGEN A --ANTAGONISTS AND INHIBITORS --AI
               2 PEPSINOGEN A --BIOSYNTHESIS --BI
              56 PEPSINOGEN A --BLOOD --BL
              12 PEPSINOGEN A --CHEMISTRY --CH
               3 PEPSINOGEN A -- DRUG EFFECTS -- DE
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10 PEPSINOGEN A --GENETICS --GE
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             В
                 13 ENZYME PRECURSORS
R6
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                 7 PEPSINOGEN A
R7
        23
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                  4 PEPSINOGEN C
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Ref
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E2
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                 PG-1
E3
        909
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E4
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E5
        11
                 PGAA
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E7
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E9
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                 PGAB
E10
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                  PGABA
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E12
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               R1-R3
S5
         133
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S6
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09774848
          99070725
   Serum
          antibodies
                       to H+,K+- ATPase, serum pepsinogen A and
Helicobacter
                 pylori in relation to gastric mucosa morphology in
patients with low or low-normal concentrations of serum cobalamins.
 Lindgren A; Burman P; Kilander AF; Nilsson O; Lindstedt G
              οf
                   Internal Medicine, Sahlgrenska University Hospital,
Goteborg, Sweden.
 European journal of gastroenterology & hepatology (ENGLAND)
                                                              Jul 1998
10 (7) p583-8, ISSN 0954-691X Journal Code: B9X
 Languages: ENGLISH
 Document type: JOURNAL ARTICLE
 JOURNAL ANNOUNCEMENT: 9904
            INDEX MEDICUS
 Subfile:
 OBJECTIVES: To compare the diagnostic performance of serum antibodies to
H+,K+- ATPase
              (EC 3.6.1.36), serum pepsinogen A (EC 3.4.23.1) and the
Schilling test in diagnosing chronic atrophic body gastritis; to study the
                             H+, K+- ATPase antibodies, serology for
interrelationships
                   between
Helicobacter
                 pylori , and gastric morphology. DESIGN: Patients with
suspected cobalamin deficiency and serum cobalamin < 200 micromol/l were
investigated using upper gastrointestinal endoscopy, the Schilling test and
serum tests for H+,K+-ATPase antibodies, pepsinogen A, and H. pylori .
SETTING: The Department of Internal Medicine, Sahlgrenska University
Hospital, Goteborg, Sweden. PATIENTS: Ninety seven consecutively referred
patients. MAIN OUTCOME MEASURES: Sensitivity and specificity of assays for
serum H+,K+- ATPase antibodies, serum pepsinogen A, and the Schilling
      RESULTS: Assays of serum antibodies to H+, K+-ATPase and of serum
              A displayed equal diagnostic sensitivity for atrophic
pepsinogen
gastritis (around 0.90 for the severe forms) and higher than that for the
Schilling test (0.65). The diagnostic specificity for pepsinogen A (1.0)
was higher than for H+,K+-ATPase antibodies (about 0.80). The prevalence
of antral gastritis and positivity for H. pylori antibodies declined with
the transition of body gastritis into severe atrophy, while the prevalence
of H+,K+- ATPase antibodies increased. CONCLUSION: Pepsinogen A is
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E9 •

E10

E11

E12

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PG100

PG101

PG11

PG10.2 PROTEIN

Enter P or PAGE for more

*preferable to serum H+,K+-ATPase antibodies in the diagnosis of gastric body mucosal atrophy. The formation of H+,K+-ATPase antibodies does not seem to be a primary event in the development of gastric body muscosal atrophy.

Tags: Human

Descriptors: Antibodies, Bacterial--Blood--BL; *Gastritis, Atrophic --Diagnosis--DI; * H(+)-K(+)-Exchanging ATPase --Bloo d--BL; * Helicobacter pylori --Immunology--IM; * Pepsinogen A--Blood--BL; *Vitamin B 12--Blood--BL; Adult; Aged; Chronic Disease; Gastric Mucosa --Immunology--IM; Gastric Mucosa--Pathology--PA; Gastritis, Atrophic --Immunology--IM; H(+)-K(+)-Exchanging ATPase --Immu nology--IM; Middle Age; Pepsinogen A--Immunology--IM; Reference Values; Schilling Test; Sensitivity and Specificity; Serologic Tests; Vitamin B 12--Immunology--IM CAS Registry No.: 0 (Antibodies, Bacterial); 68-19-9 (Vitamin B 12); 9001-10-9 (Pepsinogen A)

Enzyme No.: EC 3.6.1.36 (H(+)-K(+)-Exchanging ATPase)

10/9/2

DIALOG(R) File 155: MEDLINE(R)

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07425784 92221233

Helicobacter pylori and hypergastrinaemia during proton pump inhibitor therapy.

McColl KE; Nujumi AM; Dorrian CA; Macdonald AM; Fullarton GM; Harwood J University Dept. of Medicine and Therapeutics, Western Infirmary, Glasgow, Scotland.

Scandinavian journal of gastroenterology (NORWAY) (1992, 27 (2) p93-8

ISSN 0036-5521 Journal Code: UCS

Languages: ENGLISH

Document type: JOURNAL ARTICLE JOURNAL ANNOUNCEMENT: 9207 Subfile: INDEX MEDICUS

The rise in serum gastrin and pepsinogen I after 5 days' treatment with the proton pump inhibitor pantoprazole (40 mg/day) was examined in eight duodenal ulcer patients with Helicobacter pylori infection and compared with eight in whom it had been eradicated. Before treatment, the post-prandial serum gastrin concentrations were higher in the H. pylori positive than eradicated patients (pless than 0.05). The median rise in

-positive than -eradicated patients (p less than 0.05). The median rise in pre-prandial serum gastrin concentrations on treatment was similar in the H. pylori -positive (41%) and -eradicated patients (45%). The rise in post-prandial serum gastrin was also similar in the H. pylori -positive (81%) and -eradicated patients (69%), resulting in significantly higher gastrin concentrations during treatment in the former. The median rise in serum pepsinogen I on treatment was greater in the H. pylori -positive (114%) than in the -eradicated patients (8%), resulting in significantly higher concentrations during treatment in the former. These observations indicate that eradication of H. pylori may be a means of moderating the

indicate that eradication of H. pylori may be a means of moderating the hypergastrinaemia caused by acid-inhibitory therapy. They also indicate that H. pylori -related hypergastrinaemia is not due to an increase of the antral surface pH by the bacterium's urease activity.

Tags: Human; Male; Support, Non-U.S. Gov't

Descriptors: Adenosinetriphosphatase --Antagonists and Inhibitors--AI;
*Benzimidazoles--Pharmacology--PD; *Gastrins--Blood--BL; * Helicobacter
pylori ; *Helicobacter Infections--Blood--BL; *Pepsinogens--Blood--BL;
*Peptide Fragments--Blood--BL; *Sulfoxides--Pharmacology--PD; Benzimidazole
s--Therapeutic Use--TU; Duodenal Ulcer--Blood--BL; Duodenal Ulcer--Drug
Therapy--DT; Duodenal Ulcer--Metabolism--ME; Gastric Acidity Determination
; Gastrins--Drug Effects--DE; Helicobacter Infections--Metabolism--ME;
Pepsinogens--Drug Effects--DE; Peptide Fragments--Drug Effects--DE;
Sulfoxides--Therapeutic Use--TU

CAS Registry No.: 0 (Benzimidazoles); 0 (Gastrins); 0 (Pepsinogens); 0 (Peptide Fragments); 0 (Sulfoxides); 102625-70-7 (pantoprazole); 75903-15-0 (pepsinogen (1-12))

Enzyme No.: EC 3.6.1.3 Adenosinetriphosphatase)

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10/9/3

DIALOG(R) File 155: MEDLINE(R)

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06651765 91047803

Acid and barriers. Current research and future developments for peptic ulcer therapy.

Rademaker JW; Hunt RH

Division of Gastroenterology, McMaster University Medical Centre, Hamilton, Ontario, Canada.

Scandinavian journal of gastroenterology. Supplement (NORWAY) 1990, 175 p19-26, ISSN 0085-5928 Journal Code: UCT

Languages: ENGLISH

Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL

JOURNAL ANNOUNCEMENT: 9102 Subfile: INDEX MEDICUS

Medical therapy for peptic ulcer disease has been targeted at inhibiting acid secretion based on the belief that ulcers occur due to an imbalance between aggressive and protective factors. New antisecretory agents are unlikely to show any dramatic improvement over the success and safety of histamine H2 receptor antagonists or the recently introduced H+K+ATPase proton pump antagonist omeprazole. The development of specific muscarinic M3 and gastrin receptor antagonists will provide useful agents to suppress acid and pepsinogen secretion by alternative means and may prevent the associated hypergastrinaemia seen with anti-secretory therapy. Enhancement of mucosal defence by site protective agents will be based on a better understanding of the vascular and immune factors involved in maintaining mucosal integrity and the growth factors that regulate wound healing. Molecular techniques are likely to produce the 'model anti-ulcer' agent which will effectively inhibit acid secretion and also enhance wound healing thus providing a cure for this chronic disease. (67 Refs.)

Tags: Human

Descriptors: *Antacids--Therapeutic Use--TU; *Anti-Ulcer Agents --Therapeutic Use--TU; *Peptic Ulcer--Drug Therapy--DT; Gastric Mucosa --Physiology--PH; Helicobacter pylori; Helicobacter Infections --Complications--CO; Histamine H2 Antagonists--Therapeutic Use--TU; Intestinal Mucosa--Physiology--PH; Peptic Ulcer--Etiology--ET; Wound Healing--Physiology--PH

CAS Registry No.: 0 (Antacids); 0 (Anti-Ulcer Agents); 0 (Histamine H2 Antagonists)

10/9/4

DIALOG(R) File 155: MEDLINE(R)

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06279395 89283673

NSAIDs: new approaches to limiting gastropathy.

Zeidler H; Munzel P

Abt. Rheumatologie, Zentrum Innere Medizin und Dermatologie, Medizinische Hochschule, Hanover, West Germany.

Scandinavian journal of rheumatology. Supplement (SWEDEN) 1989, 78 pl8-23; discussion 30-2, ISSN 0301-3847 Journal Code: UD0

Languages: ENGLISH

Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL

JOURNAL ANNOUNCEMENT: 8909

Subfile: INDEX MEDICUS

An extensive literature search on non-steroidal anti-inflammatory drug (NSAID)-induced gastropathy in rheumatic conditions has been carried out. A reduced incidence of gastropathy has been observed among newly developed NSAIDs such as etodolac and the non-acidic nabumetone. An alternative prophylactic therapy to avoid NSAID-induced gastroduodenal mucosal damage which has been successfully tested in several trials is co-medication with the prostaglandin analogue misoprostol. The cytoprotective agent sucralfate also appears to be effective. Recent observations of Campylobacter pylori infections in NSAID-induced gastropathy introduces the question as to

whether simultaneous antibacterial medication should be routinely administered during NSAID therapy. At present the invasive technique of endoscopy is used to ascertain gastroduodenal mucosal damage. However, a new technique which merely requires blood sampling is being investigated. This involves measurement of serum levels of the precursor molecules for the gastric enzyme pepsin, pepsinogen I and II. In future this assay could constitute a non-invasive method for detecting gastroduodenal mucosal damage. (9 Refs.)

Tags: Human

Descriptors: *Anti-Inflammatory Agents, Non-Steroidal--Adverse Effects --AE; *Stomach Diseases--Prevention and Control--PC; Adenosinetriphosphata

Descriptors: *Anti-Inflammatory Agents, Non-Steroidal--Adverse Effects --AE; *Stomach Diseases--Prevention and Control--PC; Adenosinetriphosphata se --Antagonists and Inhibitors--AI; Anti-Inflammatory Agents, Non-Steroidal--Therapeutic Use--TU; Campylobacter Infections--Complications --CO; Campylobacter Infections--Drug Therapy--DT; Chemistry, Pharmaceutical; Drug Therapy, Combination; Pepsinogens--Blood--BL; Prostaglandins--Therapeutic Use--TU; Risk Factors; Stomach Diseases --Chemically Induced--CI; Stomach Diseases--Complications--CO; Sucralfate --Therapeutic Use--TU

CAS Registry No.: 0 (Pepsinogens); 0 (Prostaglandins); 54182-58-0 (Sucralfate)

Enzyme No.: EC 3.6.1.3 Adenosinetriphosphatase) ?logoff hold

23feb01 10:33:36 User228206 Session D1422.2 \$5.31 1.661 DialUnits File155 \$0.80 4 Type(s) in Format 9 \$0.80 4 Types

\$6.11 Estimated cost File155

\$0.35 TYMNET

\$6.46 Estimated cost this search

\$6.47 Estimated total session cost 1.835 DialUnits

Status: Signed Off. (7 minutes)

Helicobacter pylori and hypergastrinaemia during proton pump inhibitor therapy.

McColl KE; Nujumi AM; Dorrian CA; Macdonald AM; Fullarton GM; Harwood J University Dept. of Medicine and Therapeutics, Western Infirmary, Glasgow, Scotland.

Scandinavian journal of gastroenterology (NORWAY) 1992, 27 (2) p93-8, ISSN 0036-5521 Journal Code: UCS

Languages: ENGLISH

Document type: JOURNAL ARTICLE JOURNAL ANNOUNCEMENT: 9207 Subfile: INDEX MEDICUS

The rise in serum gastrin and pepsinogen I after 5 days' treatment with the proton pump inhibitor pantoprazole (40 mg/day) was examined in eight duodenal ulcer patients with Helicobacter pylori infection and compared eight in whom it had been eradicated. Before treatment, the post-prandial serum gastrin concentrations were higher in the H. pylori -positive than -eradicated patients (p less than 0.05). The median rise in pre-prandial serum gastrin concentrations on treatment was similar in the H. pylori -positive (41%) and -eradicated patients (45%). The rise in post-prandial serum gastrin was also similar in the H. pylori -positive (81%) and -eradicated patients (69%), resulting in significantly higher gastrin concentrations during treatment in the former. The median rise in serum pepsinogen I on treatment was greater in the H. pylori -positive (114%) than in the -eradicated patients (8%), resulting in significantly higher concentrations during treatment in the former. These observations indicate that eradication of H. pylori may be a means of moderating the hypergastrinaemia caused by acid-inhibitory therapy. They also indicate that H. pylori -related hypergastrinaemia is not due to an increase of the antral surface pH by the bacterium's urease activity.

Tags: Human; Male; Support, Non-U.S. Gov't

NSAIDs: new approaches to limiting gastropathy.

Zeidler H; Munzel P

Abt. Rheumatologie, Zentrum Innere Medizin und Dermatologie, Medizinische Hochschule, Hanover, West Germany.

Scandinavian journal of rheumatology. Supplement (SWEDEN) 1989, 78-p18-23; discussion 30-2, ISSN 0301-3847 Journal Code: UD0

Languages: ENGLISH

Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL

JOURNAL ANNOUNCEMENT: 8909 Subfile: INDEX MEDICUS

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Tags: Human

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